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(54) Alteration of fatty acid profiles in plants using dominant negative mutation

(57) The preparation and use of nucleic acid fragments encoding fatty acid desaturase enzymes are described. The invention permits alteration of plant lipid composition. Chimeric genes incorporating such nucleic acid fragments with suitable regulatory sequences may be used to create transgenic plants with altered levels of unsaturated fatty acids.



Descripti n

[0001] The invention relates to the preparation and use of nucleic acid fragments encoding fatty acid desaturase enzymes to modify plant lipid composition. Chimeric genes incorporating such nucleic acid fragments and suitable regulatory sequences may be used to create transgenic plants with altered levels of unsaturated fatty acids.

[0002] Plant lipids have a variety of industrial and nutritional uses and are central to plant membrane function and climatic adaptation. These lipids represent a vast array of chemical structures, and these structures determine the physiological and industrial properties of the lipid. Many of these structures result either directly or indirectly from metabolic processes that alter the degree of unsaturation of the lipid. Different metabolic regimes in different plants produce these altered lipids, and either domestication of exotic plant species or modification of agronomically adapted species is usually required to economically produce large amounts of the desired lipid.

[0003] Plant lipids find their major use as edible oils in the form of triacylglycerols. The specific performance and health attributes of edible oils are determined largely by their fatty acid composition. Most vegetable oils derived from commercial plant varieties are composed primarily of palmitic (16:0), stearic (18:0), oleic (18:1), linoleic (18:2) and linolenic (18:3) acids. Palmitic and stearic acids are, respectively, 16- and 18-carbon-long, saturated fatty acids. Oleic, linoleic, and linolenic acids are 18-carbon-long, unsaturated fatty acids containing one, two, and three double bonds, respectively. Oleic acid is referred to as a mono-unsaturated fatty acid, while linoleic and linolenic acids are referred to as poly-unsaturated fatty acids.

[0004] Many recent research efforts have examined the role that saturated and unsaturated fatty acids play in reducing the risk of coronary heart disease. In the past, it was believed that mono-unsaturates, in contrast to saturates and poly-unsaturates, had no effect on serum cholesterol and coronary heart disease risk. Several recent human clinical studies suggest that diets high in mono-unsaturated fat and low in saturated fat may reduce the "bad" (low-density lipoprotein) cholesterol while maintaining the "good" (high-density lipoprotein) cholesterol (Mattson et al., *Journal of Lipid Research* (1985) 26:194-202).

[0005] A vegetable oil low in total saturates and high in mono-unsaturates would provide significant health benefits to consumers as well as economic benefits to oil processors. For specialized uses, high levels of poly-unsaturates can be desirable. Linoleate and linolenate are essential fatty acids in human diets, and an edible oil high in these fatty acids can be used for nutritional supplements, for example in baby foods.

[0006] The biosynthesis of the major plant lipids has been the focus of much research (Browse et al., *Ann. Rev. Plant Physiol. Mol. Biol.* (1991) 42:467-506). These studies show that, with the notable exception of the soluble stearoylacyl carrier protein desaturase, the controlling steps in the production of unsaturated fatty acids are largely catalyzed by membrane-associated fatty acid desaturases. Desaturation reactions occur in plastids and in the endoplasmic reticulum using a variety of substrates including galactolipids, sulfolipids, and phospholipids. Genetic and physiological analyses of <u>Arabidopsis thaliana</u> nuclear mutants defective in various fatty acid desaturation reactions indicates that most of these reactions are catalyzed by enzymes encoded at single genetic loci in the plant. These investigations have demonstrated the role of delta-12 desaturase and delta-15 desaturase activities in the production of linoleate and linolenate from 2-oleoyl-phosphatidylcholine and 2-linoleoyl-phosphatidylcholine, respectively (Wang et al., *Plant Physiol. Biochem.* (1988) 26:777-792). Thus, modification of the activities of these enzymes represents an attractive target for altering the levels of lipid unsaturation by genetic engineering.

[0007] The cloning and characterization of wild-type delta-12 fatty acid desaturases has been reported (Okuley, et al., Plant Cell (1994) 6:147-158). However, there are no teachings concerning plants having seed-specific expression of mutant delta-12 or delta-15 fatty acid desaturase gene products. Furthermore, no methods have been described for altering the fatty acid composition of plants using nucleic acid constructs expressing a mutant delta-12 or a mutant delta-15 fatty acid desaturase.

[0008] Applicants have discovered a means to control the nature and levels of unsaturated fatty acids in plants. Nucleic acid fragments from cDNAs or genes encoding mutant fatty acid desaturases are used to create chimeric genes. The chimeric genes may be used to transform various plants to modify the fatty acid composition of the plant or the oil produced by the plant. The invention comprises nucleic acid constructs containing mutant microsomal delta-12 or mutant microsomal delta-15 fatty acid desaturase coding sequences, which are operably linked in sense orientation to at least one regulatory sequence. Such a construct is effective for altering fatty acid composition of seeds when the construct is introduced into a plant. In one embodiment, a mutant coding sequence for a delta-12 fatty acid desaturase comprises the mutation in the sequence of SEQ ID NO:3 or SEQ ID NO:8.

[0009] Thus viewed from one aspect the present invention provides a method for altering fatty acid composition in plant seeds, comprising the steps of:

a) introducing a recombinant nucleic acid construct into a plant, said construct comprising at least one seed-specific regulatory sequence operably linked in sense orientation to a mutant delta-12 or delta-15 fatty acid desaturase gene encoding a protein having a mutation in a His-(Asp/Glu)-Cys-(Gly/Ala)-His amino acid region;

- b) obtaining progeny from said plant, said progeny producing seeds having said altered fatty acid composition; and c) producing seeds having said altered fatty acid composition.
- [0010] Constructs containing both mutant delta-12 and delta-15 fatty acid desaturase genes may also be used in this method.
- [0011] Depending on the mutant that is introduced, alterations in the linoleic acid content (for example from about 1.0% to about 10%, based on total fatty acid composition), oleic acid content (for example from about 69% to about 90%) or α -linolenic acid (for example from about 1.0% to about 10%) may be achieved.
- [0012] Preferably the mutant desaturase gene used in the construct comprises the full-length coding sequence.
- [0013] Preferably said mutant gene is derived from a Brassica napus plant.
 - [0014] Constructs for use in the above method, and mutant fatty acid desaturase proteins form further features of the invention.
 - [0015] The invention further comprises seeds, which may be in non-viable form, such as crushed or milled, the meal derived therefrom, plants, plant lines and plant cells having a recombinant nucleic acid construct containing at least one regulatory sequence linked in sense orientation to a mutant delta-12 or mutant delta-15 fatty acid desaturase. The mutant chimeric gene preferentially is expressed in seeds and results in an altered fatty acid composition in seeds of such plants. A plant expressing a mutant delta-12 desaturase gene preferably has a reduced level of linoleic acid in seeds. A plant expressing a mutant delta-15 desaturase gene preferably has a reduced level of α -linolenic acid in seeds. If desired, a plant of the invention may express both a mutant delta-12 and a mutant delta-15 fatty acid desaturase, resulting in the reduction of both linoleic acid and α -linolenic acid in seeds.
 - [0016] Yet another embodiment of the invention involves a method of producing seed oil containing altered levels of unsaturated fatty acids comprising: (a) transforming a plant cell with a chimeric gene described above; (b) growing sexually mature plants from the transformed plant cells of step (a); (c) screening progeny seeds from the sexually mature plants of step (b) for the desired levels of unsaturated fatty acids, and (d) processing the progeny seed of step (c) to obtain seed oil containing altered levels of the unsaturated fatty acids. Preferred plant cells and oils are derived from soybean, rapeseed, sunflower, cotton, cocoa, peanut, safflower, coconut, flax, oil palm, and corn. Preferred methods of transforming such plant cells would include the use of Ti and Ri plasmids of <u>Agrobacterium</u>, electroporation, and high-velocity ballistic bombardment.
- [0017] Yet another aspect of the invention involves a method of producing seeds having altered fatty acid composition. The method comprises the step of introducing a recombinant nucleic acid construct into a plant (i.e., transforming a plant). The construct comprises one or more seed-specific regulatory sequences operably linked in sense orientation to a mutant delta-12 fatty acid desaturase gene or a mutant delta-15 fatty acid desaturase gene. After obtaining transgenic progeny, those transformed plants that produce seeds having an altered fatty acid composition are identified. Suitable plants for transformation include, for example, soybean, rapeseed, sunflower, safflower, castor bean and corn. Suitable methods of transforming such plants include, for example, Agrobacterium-mediated methods, electroporation, and microprojectile bombardment.
 - [0018] The invention also is embodied in a method of RFLP breeding to obtain altered levels of oleic acids in the seed oil of oil producing plant species. This method involves (a) making a cross between two varieties of oil producing plant species differing in the oleic acid trait; (b) making a Southern blot of restriction enzyme digested genomic DNA isolated from several progeny plants resulting from the cross; and (c) hybridizing the Southern blot with the radiolabelled nucleic acid fragments encoding the mutant fatty acid desaturases or desaturase-related enzymes.
 - [0019] The invention is also embodied in a method of RFLP mapping that uses the isolated mutant microsomal delta-12 desaturase cDNA or related genomic fragments described herein.
- [0020] Another embodiment of the instant invention is a method of genotyping plants containing either a mutant or wild-type form of the delta-12 desaturase gene by PCR amplification of genomic DNA using gene-specific primers. This method is capable of discriminating genes that differ by only one or a few nucleotides, thus affording a means for detecting plants containing the mutant delta-12 desaturase.
- [0021] Another aspect of the invention comprises vegetable oil extracted from seeds of plants disclosed herein. Such a vegetable oil contains an altered fatty acid composition, e.g., a decreased level of α -linolenic acid, a decreased level of linoleic acid, or an increased level of oleic acid, based on total fatty acid composition.
- [0022] The invention can be more fully understood from the following detailed description and the Sequence Descriptions which form a part of this application. The Sequence Descriptions contain the three letter codes for amino acids as defined in WIPO STANDARD ST.23.
- [0023] SEQ ID NO:1 shows the 5' to 3' nucleotide sequence of 1464 base pairs of the <u>Brassica napus</u> cDNA which encodes the wild type D form of microsomal delta-12 desaturase in plasmid pCF2-165d.
- [0024] SEQ ID NO:2 is the 384 amino acid protein sequence deduced from the open reading frame in SEQ ID NO:1. [0025] SEQ ID NO:3 shows the 5' to 3' cDNA nucleotide sequence of a mutant D form of microsomal delta-12 fatty acid desaturase from Brassica napus IMC129. Nucleotides 1-3 are the initiation codon and nucleotides 1153-1155 are

the termination codon.

[0026] SEQ ID NO:4 is the 384 amino acid protein sequence deduced from the open reading frame of SEQ ID NO:3. [0027] SEQ ID NO:5 shows the 5' to 3' cDNA nucleotide sequence of the wild-type F form of microsomal delta-12 fatty acid desaturase in <u>Brassica napus</u>. Nucleotides 1-3 are the initiation codon and nucleotides 1153-1155 are the termination codon.

[0028] SEQ ID NO:6 is the 384 amino acid protein sequence deduced from the open reading frame of SEQ ID NO:5. [0029] SEQ ID NO:7 shows the 5' to 3' cDNA nucleotide sequence of a mutant F form of microsomal delta-12 fatty acid desaturase from <u>Brassica napus</u> IMC Q508. Nucleotides 1-3 are the initiation codon and nucleotides 1153-1155 are the termination codon.

[0030] SEQ ID NO:8 is the 384 amino acid protein sequence deduced from the open reading frame of SEQ ID NO:7.

[0031] SEQ ID NO:9 is the upstream (5') primer used for isolation of the D form of microsomal delta-12 fatty acid desaturase gene from <u>Brassica napus</u>.

[0032] SEQ ID NO:10 is the downstream (3') primer used for isolation of the D form of microsomal delta-12 fatty acid desaturase gene from <u>Brassica napus</u>.

[0033] SEQ ID NO:11 is the upstream (5') primer used for isolation of the F form of microsomal delta-12 fatty acid desaturase gene in Brassica napus.

[0034] SEQ ID NO:12 is the downstream (3') primer used for isolation of the F form of microsomal delta-12 fatty acid desaturase gene in <u>Brassica napus</u>.

[0035] SEQ ID NO:13 is the upstream (5') primer used for gene-specific detection of the wild type D form of microsomal delta-12 fatty acid desaturase gene in <u>Brassica napus</u>.

[0036] SEQ ID NO:14 is the upstream (5') primer used for gene-specific detection of the mutant D form of microsomal delta-12 fatty acid desaturase gene in <u>Brassica napus</u>.

[0037] SEQ ID NO:15 is the modified upstream (5) primer used for gene-specific detection of the wild type D form of microsomal delta-12 fatty acid desaturase gene in <u>Brassica napus</u>.

[0038] SEQ ID NO:16 is the modified upstream (5') primer used for gene-specific detection of the mutant D form of microsomal delta-12 fatty acid desaturase gene in <u>Brassica napus</u>.

[0039] The Examples herein are given by way of illustration only in which the Figures referred to are as follows:

Figure 1 is a schematic drawing of plasmid pZPhMcFd2, showing restriction sites and relative position and orientation of the bean phaseolin promoter (5' Phas), the IMC129 mutant microsomal delta-12 fatty acid desaturase D form coding sequence (MCFd2) and the bean phaseolin 3' untranslated region (3' Phas).

Figure 2 is a schematic drawing of plasmid pIMC127, showing restriction sites and the relative positions and orientation of the napin promoter (5' nap), the wild-type microsomal delta-12 fatty acid desaturase D form coding sequence (CanFd2) and the napin 3' untranslated region (3' Nap).

Figure 3 shows the frequency distribution of seed oil linoleic acid (C18:2) content in transgenic Brassica T2 populations transformed with either a napin promoter linked in sense orientation to a wild-type microsomal delta-12 fatty acid desaturase D form coding sequence (WS127) or a phaseolin promoter linked to a mutant delta-12 fatty acid desaturase D form (WS201).

Figure 4 shows the frequency distribution of seed oil linoleic acid content in transgenic Brassica T2 populations transformed with either a napin promoter linked in sense orientation to a mutant F form (WS140) delta-12 fatty acid desaturase coding sequence or a cruciferin promoter linked to a wild-type delta-12 fatty acid desaturase D form (WS135).

[0040] Nucleic acid fragments have been isolated that encode mutant plant fatty acid desaturases and that are useful in modifying fatty acid composition in oil-producing species by genetic transformation.

[0041] Transfer of the nucleic acid fragments of the invention or a part thereof, along with suitable regulatory sequences that direct the transcription of their mRNA, into plants may result in production of decreased levels of unsaturated fatty acids in cellular lipids, including triacylglycerols.

[0042] The nucleic acid fragments of the invention can also be used as DNA diagnostic markers in plant genetic mapping and plant breeding programs.

[0043] The nucleic acid fragments of the invention or oligomers derived therefrom can also be used to isolate other related fatty acid desaturase genes using DNA, RNA, or a library of cloned nucleotide sequences from the same or different species by well known sequence-dependent protocols, including, for example, methods of nucleic acid hybridization and amplification by the polymerase chain reaction.

[0044] In the context of this disclosure, a number of terms shall be used. Fatty acids are specified by the number of carbon atoms and the number and position of the double bond: the numbers before and after the colon refer to the chain length and the number of double bonds, respectively. The number following the fatty acid designation indicates the position of the double bond from the carboxyl end of the fatty acid with the "c" affix for the cis-configuration of the double

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bond. For example, palmitic acid (16:0), stearic acid (18:0), oleic acid (18:1,9c), petroselinic acid (18:1, 6c), linoleic acid (18:2,9c,12c), γ -linolenic acid (18:3, 6c,9c,12c) and α -linolenic acid (18:3, 9c,12c,15c). Unless otherwise specified 18:1, 18:2 and 18:3 refer to oleic, linoleic and linolenic fatty acids. The term "fatty acid desaturase" used herein refers to an enzyme which catalyzes the breakage of a carbon-hydrogen bond and the introduction of a carbon-carbon double bond into a fatty acid molecule. The fatty acid may be free or esterified to another molecule including, but not limited to, acylcarrier protein, coenzyme A, sterols and the glycerol moiety of glycerolipids. The term "glycerolipid desaturases" used herein refers to a subset of the fatty acid desaturases that act on fatty acyl moieties esterified to a glycerol backbone. "Delta-12 desaturase" refers to a fatty acid desaturase that catalyzes the formation of a double bond between carbon positions 6 and 7 (numbered from the methyl end), (i.e., those that correspond to carbon positions 12 and 13 (numbered from the carbonyl carbon) of an 18 carbon-long fatty acyl chain. "Delta-15 desaturase" refers to a fatty acid desaturase that catalyzes the formation of a double bond between carbon positions 3 and 4 (numbered from the methyl end), (i.e., those that correspond to carbon positions 15 and 16 (numbered from the carbonyl carbon) of an 18 carbon-long fatty acyl chain. Examples of fatty acid desaturases include, but are not limited to, the microsomal delta-12 and delta-15 desaturases that act on phosphatidylcholine lipid substrates; the chloroplastic or plastid delta-12 and delta-15 desaturases that act on phosphatidyl glycerol and galactolipids; and other desaturases that act on such fatty acid substrates such as phospholipids, galactolipids, and sulfolipids. "Microsomal desaturase" refers to the cytoplasmic location of the enzyme, while "chloroplast desaturase" and "plastid desaturase" refer to the plastid location of the enzyme. These fatty acid desaturases may be found in a variety of organisms including, but not limited to, higher plants, diatoms, and various eukaryotic and prokaryotic microorganisms such as fungi and photosynthetic bacteria and algae. The term "homologous fatty acid desaturases" refers to fatty acid desaturases that catalyze the same desaturation on the same lipid substrate. Thus, microsomal delta-15 desaturases, even from different plant species, are homologous fatty acid desaturases. The term "heterologous fatty acid desaturases" refers to fatty acid desaturases that catalyze desaturations at different positions and/or on different lipid substrates. Thus, for example, microsomal delta-12 and delta-15 desaturases, which act on phosphatidylcholine lipids, are heterologous fatty acid desaturases, even when from the same plant. Similarly, microsomal delta-15 desaturase, which acts on phosphatidylcholine lipids, and chloroplast delta-15 desaturase, which acts on galactolipids, are heterologous fatty acid desaturases, even when from the same plant. It should be noted that these fatty acid desaturases have never been isolated and characterized as proteins. Accordingly, the terms such as "delta-12 desaturase" and "delta-15 desaturase" are used as a convenience to describe the proteins encoded by nucleic acid fragments that have been isolated based on the phenotypic effects caused by their disruption. They do not imply any catalytic mechanism. For example, delta-12 desaturase refers to the enzyme that catalyzes the formation of a double bond between carbons 12 and 13 of an 18 carbon fatty acid irrespective of whether it "counts" the carbons from the methyl, carboxyl end, or the first double bond.

[0045] Desaturases which are mutated according to the invention include both naturally-occurring enzymes (produced either naturally or synthetically) and functionally-equivalent derivatives and variants thereof. Such derivatives and variants, which may be related to or derived from native proteins include proteins with variation in the amino acid sequence and/or particular residues in the sequence without significantly affecting the desaturase functionality of the protein (ie. functionally equivalent). Such functionally-equivalent variants or derivatives include natural biological variations (e.g. allelic variants or geographical variants within a species, or variants, such as analogues from different species or genera) and derivatives prepared using known techniques. For example, functionally-equivalent proteins may be prepared by chemical peptide synthesis or in recombinant form using known techniques of site-directed mutagenesis, random mutagenesis, or enzymatic cleavage and/or ligation of nucleic acids. Thus for example, alteration by single or multiple substitution, modification, deletion and/or addition of the amino acid sequence is contemplated.

[0046] Appropriate alterations which retain the protein's functionality will be clear to the person skilled in the art. Such alterations include for example conservative substitution of residues, such as for example replacement of a hydrophobic amino acid with a different hydrophobic amino acid, for example glycine, valine, leucine or isoleucine. Similarly, negatively charged residues may be replaced for example in the case of glutamic acid with aspartic acid, or vice versa, and in the case of positively charged residues arginine may be replaced with lysine or vice versa.

[0047] In certain cases modification of amino acids is appropriate and in these case this may be performed pre- or post-translationally. Thus for example, non-natural amino acids may be used in synthesis or modified (e.g. methylation) after translation. Modification includes for example glycosylation, deglycosylation, phosphorylation etc. It will be appreciated that modification of the amino acid sequence pre- or post-translationally will be of limited use for methods of altering fatty acid compositions as described herein which relies on expression of mutant genes. The present invention is however also concerned with novel mutant desaturases themselves which may be modified as described above.

[0048] Preferably the derivatives or variants defined above are substantially similar to desaturases known in the art or described in the present application, ie. the alterations ar of a relatively minor nature involving for example conservative substitution and/or single additions/deletions and/or modification of the amino acids. When reference is made to functionally-equivalent variants of mutant desaturases according to the invention, the alterations described above apply to the non-mutated portions of the sequence.

[0049] Functionally-equivalent protein "fragments" as used herein refers to portions of the protein, e.g. desaturase which contains at least the domain(s) responsible for the enzymatic activity of the enzyme. When reference is made to fragments of mutant desaturases, the fragment corresponds to the functionally-equivalent fragment of the desaturase as defined above, which has been mutated according to the invention.

[0050] The term "nucleic acid" refers to a large molecule which can be single-stranded or double-stranded, composed of monomers (nucleotides) containing a sugar, a phosphate and either a purine or pyrimidine. A "nucleic acid fragment" is a fraction of a given nucleic acid molecule. In higher plants, deoxyribonucleic acid (DNA) is the genetic material while ribonucleic acid (RNA) is involved in the transfer of the information in DNA into proteins. A "genome" is the entire body of genetic material contained in each cell of an organism. The term "nucleotide sequence" refers to the sequence of DNA or RNA polymers, which can be single- or double-stranded, optionally containing synthetic, non-natural or altered nucleotide bases capable of incorporation into DNA or RNA polymers. The term "oligomer" refers to short nucleotide sequences, usually up to 100 bases long.

[0051] From time to time, the term "FAD2" may be used herein as a shorthand notation for a nucleotide sequence encoding a wild type microsomal delta-12 fatty acid desaturase enzyme, and the term "fad2" may be used herein as a shorthand notation for a nucleotide sequence encoding a mutant form of a microsomal delta-12 fatty acid desaturase enzyme.

[0052] As used herein, the term "homologous to" refers to the relatedness between the nucleotide sequence of two nucleic acid molecules or between the amino acid sequences of two protein molecules. Estimates of such homology are provided by either DNA-DNA or DNA-RNA hybridization under conditions of stringency as is well understood by "those skilled in the art (Hames and Higgins, Eds. (1985) Nucleic Acid Hybridisation, IRL Press, Oxford, U.K.); or by the comparison of sequence similarity between two nucleic acids or proteins, such as by the method of Needleman et al. (J. Mol. Biol. (1970) 48:443-453). As used herein, "essentially similar" refers to DNA sequences that may involve base changes that do not cause a change in the encoded amino acid, or which involve base changes which may alter one or more amino acids, but do not affect the functional properties of the protein encoded by the DNA sequence. It is therefore understood that the invention encompasses more than the specific exemplary sequences. Modifications to the sequence, such as deletions, insertions, or substitutions in the sequence which produce silent changes that do not substantially affect the functional properties of the resulting protein molecule are also contemplated.

[0053] Appropriate nucleic acid sequence variants encoding functionally-equivalent variants of the desaturases or mutant desaturases of the invention include for example substantially homologous sequences, for example displaying at least 50 or 60%, preferably at least 70, 80 or 90% sequence homology. Hybridizing nucleic acid sequences also constitute appropriate variants, for example those which bind under non-stringent conditions (6xSC/50% formamide at room temperature) and washed under conditions of low stringency (2XSSC, room temperature, more preferably 2XSSC, 42°C) or conditions of higher stringency e.g. 2XSSC, 65°C (where SSC = 0.15M NaCl, 0.015M sodium citrate, pH 7.2), as well as those which, but for the degeneracy of the code, would hybridize under the above-mentioned conditions

[0054] Thus, in particular for example, alteration in the gene sequence which reflect the degeneracy of the genetic code, or which results in the production of a chemically equivalent amino acid at a given site, are contemplated; thus, a codon for the amino acid alanine, a hydrophobic amino acid, may be substituted by a codon encoding another hydrophobic amino acid residue such as glycine, valine, leucine, or isoleucine. Similarly, changes which result in substitution of one negatively charged residue for another, such as aspartic acid for glutamic acid, or one positively charged residue for another, such as lysine for arginine, can also be expected to produce a biologically equivalent product. Nucleotide changes which result in alteration of the N-terminal and C-terminal portions of the protein molecule would also not be expected to alter the activity of the protein. Each of the proposed modifications is well within the routine skill in the art, as is determination of retention of biological activity of the encoded products.

[0055] "Gene" refers to a nucleic acid fragment that encodes a specific protein, including regulatory sequences preceding (5' non-coding) and following (3' non-coding) the coding region. "Fatty acid desaturase gene" refers to a nucleic acid fragment that encodes a protein with fatty acid desaturase activity. "Native" gene refers to an isolated gene with its own regulatory sequences as found in nature. "Chimeric gene" refers to a gene that comprises heterogeneous regulatory and coding sequences not found in nature. "Endogenous" gene refers to the native gene normally found in its natural location in the genome and is not isolated. A "foreign" gene refers to a gene not normally found in the host organism but that is introduced by gene transfer. "Pseudo-gene" refers to a genomic nucleotide sequence that does not encode a functional enzyme. "Mutant gene" refers to a gene comprising one or more nucleotides that have been altered when compared to the wild-type nucleotide sequence, resulting in a change to the amino acid sequence and functional properties of the encoded protein. The mutant genes may be derived from wild-type genes, such as from <u>Brassica napus</u>. However, as indicated herein, gen sequences which may be mutated include any sequences which encode proteins of

the desired functionality before mutation. Thus, a "mutation" is a change which occurs in a gene to give rise to an altered form of the gene which differs from the wild-type gene. As those skilled in the art will appreciate such changes can occur due to a deletion, addition, substitution or combination thereof of the nucleic acid sequence.

[0056] Mutant desaturase proteins encoded by said mutant genes exhibit altered functional properties as described herein. Mutations may be made in the proteins at any appropriate site or sites within the protein. Examples of appropriate sites which may be mutated include one or more amino acids in regions which are conserved between different genera and/or species. Examples of appropriate conserved regions in this connection may be found in Table 1. Particularly preferred are mutations in the consensus sequence of delta 12-desaturases and delta-15 desaturases such as in the His-(Asp/Glu)-Cys-(Gly/Ala)-His region. Preferred mutations include, but are not limited to non-conservative substitutions of one or more amino acids. For example, as described herein, a glutamic acid residue may be replaced with a lysine residue and/or a leucine residue may be replaced with a histidine residue.

[0057] "Coding sequence" refers to a DNA sequence that codes for a specific protein and excludes the non-coding sequences. It may constitute an "uninterrupted coding sequence", i.e., lacking an intron or it may include one or more introns bounded by appropriate splice junctions. An "intron" is a nucleotide sequence that is transcribed into a primary transcript but that is removed through cleavage and re-ligation of the RNA within the cell to create a mature mRNA that can be translated into a protein.

[0058] "Initiation codon" and "termination codon" refer to a unit of three adjacent nucleotides in a coding sequence that specifies initiation and chain termination, respectively, of protein synthesis (mRNA translation). "Open reading frame" refers to the coding sequence uninterrupted by introns between initiation and termination codons that encodes an amino acid sequence.

[0059] "RNA transcript" refers to the product resulting from RNA polymerase-catalyzed transcription of a DNA sequence. When the RNA transcript is a perfect complementary copy of the DNA sequence, it is referred to as the primary transcript or it may be a RNA sequence derived from posttranscriptional processing of the primary transcript and is referred to as the mature RNA. "Messenger RNA (mRNA)" refers to the RNA that is without introns and that can be translated into protein by the cell. "cDNA" refers to a double-stranded DNA that is complementary to and derived from mRNA. "Sense" RNA refers to RNA transcript that includes the mRNA. "Antisense RNA" refers to a RNA transcript that is complementary to all or part of a target primary transcript or mRNA and may block the expression of a target gene by interfering with the processing, transport and/or translation of its primary transcript or mRNA. The complementarity of an antisense RNA may be with any part of the specific gene transcript, i.e., at the 5' non-coding sequence, 3' non-coding sequence, introns, or the coding sequence. In addition, as used herein, antisense RNA may contain regions of ribozyme sequences that increase the efficacy of antisense RNA to block gene expression. "Ribozyme" refers to a catalytic RNA and includes sequence-specific endoribonucleases.

[0060] As used herein, "suitable regulatory sequences" refer to nucleotide sequences in native or chimeric genes that are located upstream (5'), within, and/or downstream (3') to the nucleic acid fragments of the invention, which control the expression of the nucleic acid fragments of the invention. The term "expression", as used herein, refers to the transcription and stable accumulation of the sense (mRNA) derived from the nucleic acid fragment(s) of the invention that, in conjunction with the protein apparatus of the cell, results in altered levels of the fatty acid desaturase(s). Expression or overexpression of the gene involves transcription of the gene and translation of the mRNA into precursor or mature fatty acid desaturase proteins. "Altered levels" refers to the production of gene product(s) in transgenic organisms in amounts or proportions that differ from that of normal or non-transformed organisms.

[0061] "Promoter" refers to a DNA sequence in a gene, usually upstream (5') to its coding sequence, which controls the expression of the coding sequence by providing the recognition for RNA polymerase and other factors required for proper transcription. Promoters may also contain DNA sequences that are involved in the binding of protein factors which control the effectiveness of transcription initiation in response to physiological or developmental conditions. It may also contain enhancer elements. An "enhancer" is a DNA sequence which can stimulate promoter activity. It may be an innate element of the promoter or a heterologous element inserted to enhance the level and/or tissue-specificity of a promoter. "Constitutive promoters" refers to those that direct gene expression in all tissues and at all times. "Tissue-specific" or "development-specific" promoters as referred to herein are those that direct gene expression almost exclusively in specific tissues, such as leaves or seeds, or at specific development stages in a tissue, such as in early or late embryogenesis, respectively.

[0062] The "3' non-coding sequences" refers to the DNA sequence portion of a gene that contains a polyadenylation signal and any other regulatory signal capable of affecting mRNA processing or gene expression. The polyadenylation signal is usually characterized by affecting the addition of polyadenylic acid tracts to the 3' end of the mRNA precursor. [0063] "Transformation" herein refers to the transfer of a foreign gene into the genome of a host organism and its genetically stable inheritance. "Restriction fragment length polymorphism" (RFLP) refers to different sized restriction fragment lengths due to altered nucleotide sequences in or around variant forms of genes. "Molecular breeding" refers to the use of DNA-based diagnostics, such as RFLP, RAPDs, and PCR in breeding. "Fertile" refers to plants that are able to propagate sexually.

[0064] "Plants" refer to photosynthetic organisms, both eukaryotic and prokaryotic, whereas the term "Higher plants" refers to eukaryotic plants. "Oil-producing species" herein refers to plant species which produce and store triacylglycerol in specific organs, primarily in seeds. Such species include soybean (Glycine max), rapeseed and canola (includ-



ing <u>Brassica napus</u>, <u>B. campestris</u>), sunflower (<u>Helianthus annus</u>), cotton (<u>Gossypium hirsutum</u>), corn (<u>Zea mays</u>), cocoa (<u>Theobrama cacao</u>), safflower (<u>Carthamus tinctorius</u>), oil palm (<u>Elaeis guineensis</u>), coconut palm (<u>Cocos nucitera</u>), flax (<u>Linum usitatissimum</u>), (castor (<u>Ricinus commumis</u>)) and peanut (<u>Arachis hypogaea</u>). The group also includes non-agronomic species which are useful in developing appropriate expression vectors such as tobacco, rapid cycling <u>Brassica</u> species, and <u>Arabidopsis thaliana</u>, and wild species which may be a source of unique fatty acids.

[0065] "Progeny" includes descendants of a particular plant or plant line, e.g., seeds and plants of F1, F2, F3, and subsequent generations, or seeds and plants of backcrossed populations BC1, BC2, BC3 and subsequent generations.

[0066] "Sequence-dependent protocols" refer to techniques that rely on a nucleotide sequence for their utility. Examples of sequence-dependent protocols include, but are not limited to, the methods of nucleic acid and oligomer hybridization and methods of DNA and RNA amplification such as are exemplified in various uses of the polymerase chain reaction (PCR).

[0067] Various solutions used in the experimental manipulations are referred to by their common names such as "SSC", "SSPE", "Denhardt's solution", etc. The composition of these solutions may be found by reference to Appendix B of Sambrook, et al. (Molecular Cloning, A Laboratory Manual, 2nd ed. (1989), Cold Spring Harbor Laboratory Press).
[0068] WO 94/11516, incorporated herein by reference, describes the isolation and characterization of cDNAs encoding wild-type microsomal delta-12 fatty acid desaturases from a number of plant species, including <u>Arabidopsis thaliana</u>, <u>Brassica napus</u>, <u>Glycine max</u>, <u>Zea mays</u> and Castor bean. Moreover, that application demonstrates successful alteration of fatty acid content of oils from seeds obtained from transgenic plants expressing sense or antisense mRNAs encoding microsomal delta-12 fatty acid desaturases.

[0069] Alignments of protein sequences of plant microsomal fatty acid delta-12 desaturases and plant delta-15 desaturases [microsomal and plastid delta-15 desaturases from <u>Arabidopsis</u> and <u>Brassica napus</u>, WO 93/11245] allows identification of amino acid sequences conserved between the different desaturases (Table 1).

TABLE 1

Region .	Conserved AA Positions in SEQ ID NO:2 of WO94/11516	Consensus Conserved AA Sequence in Δ ¹² Desaturases	Consensus Conserved AA Sequence in Δ ¹⁵ Desaturases	Consensus AA Sequence
Α	39-44	AIPPHC	AIPKHC	AIP(P/K)HC
В	86-90	WP(L/I)YW	WPLYW	WP(L/I)YW
С	104-109	AHECGH	GHDCGH	(A/G)H(D/E)CGH
D	130-134	LLVPY	LVPY	(L/I)LVPY
Ε .	137-142	WKYSHR	WRISHR	W(K/R)(Y/I)SHR
F	140-145	SHRRHH	SHRTHH	SHR(R/T)HH
G	269-274	ITYLQ	VTYLH	(I/V)TYL(Q/H)
Н	279-282	LPHY	LPWY	LP(H/W)Y
1	289-294	WL(R/K)GAL	YLRGGL	(W/Y)L(R/K)G(A/G
J	296-302	TVDRDYG	TLDRDYG	T(V/L)DRDYG
К	314-321	THVAHHLF	THVIHHLF	THV(A/I)HHLF
L .	318-327	HHLFSTMPHY	HHLFPQIPHY	HHLF(S/P) (T/Q)(I/M)PHY

[0070] Table 1 shows twelve regions of conserved amino acid sequences, designated A-L (column 1), whose positions in SEQ ID NO:2 of WO94/11516 are shown in column 2. The consensus sequences for these regions in plant delta-12 fatty acid desaturases and plant delta-15 fatty acid desaturases are shown in columns 3 and 4, respectively; amino acids are shown by standard abbreviations, the underlined amino acids are conserved between the delta-12 and the delta-15 desaturases, and amino acids in brackets represent substitutions found at that position. The consensus sequence of these regions are shown in column 5. These short conserved amino acids and their relative positions further confirm that the isolated isolated cDNAs encode a fatty acid desaturase.

[0071] In one embodiment, transgenic plants, plant lines and plant cells according to the invention contain an introduced nucleic acid construct that comprises at least a portion of a mutant delta-12 or delta-15 desaturase coding sequence. Surprisingly, a construct comprising a mutant delta-12 desaturase or delta-15 desaturase coding sequence, operably linked in sense orientation to one or more regulatory sequences, has been found to inhibit the corresponding endogenous fatty acid desaturase activity in plants which contain such a construct. This phenomenon has been termed dominant negative suppression.

[0072] The basis for the phenomenon of dominant negative suppression is not understood. One possible explanation is that the delta-12 desaturase gene product exists as a dimer *in vivo*. If so, a dimer consisting of the mutant gene product and the wild-type gene product may be non-functional. Regardless of the actual mechanism by which dominant negative suppression operates, the observation that transformation of plants with a mutant delta-12 desaturase gene results in a large proportion of the transgenic progeny having endogenous wild-type enzyme activity inhibited by expression of the mutant gene can be used to advantage. For example, the phenomenon of dominant negative suppression can be used to alter plant desaturase enzyme activity in a tissue-specific manner. The phenomenon may also allow transformation experiments to be carried out in which a higher proportion of the resulting transgenic plants have a desired altered fatty acid profile and allow transgenic plants having desired fatty acid profiles to be more readily obtained.

[0073] Preferred constructs contain, in addition, at least one regulatory sequence operably linked in the sense orientation to the mutant coding sequence. Regulatory sequences typically do not themselves code for a gene product. Instead, regulatory sequences affect the expression level of the mutant coding sequence.

[0074] In preferred embodiments, regulatory sequences for dominant negative suppression are tissue-specific, i.e., the mutant desaturase gene product is preferentially expressed in certain plant tissues and expressed at low levels or not at all in the remaining tissues of the plant. Suitable tissue-specific regulatory sequences include those that permit expression preferentially in developing seeds. Seed-specific regulatory sequences preferably stimulate or induce levels of mutant desaturase gene product expression that coincide with the levels of wild-type desaturase gene product expression.

[0075] Dominant negative suppression plants according to the invention preferably yield seeds containing an altered fatty acid profile. For example, oil obtained from seeds of such plants may have from about 69% to about 90% oleic acid, based on the total fatty acid composition of the seed. Such oil preferably has from about 74% to about 90% oleic acid, more preferably from about 80% to about 90% oleic acid. In some embodiments, oil extracted from seeds produced by plants of the invention may have from about 3% to about 5% saturated fatty acids, based on total fatty acid composition of the seeds. In some embodiments, oil extracted from seeds of the invention may have from about 1% to about 10% α-linolenic acid.

[0076] After a recombinant nucleic acid construct, comprising a mutant microsomal delta-12 fatty acid desaturase coding sequence operably linked in the sense orientation to one or more regulatory sequences, is introduced into a plant, seeds of transgenic plants are grown and either selfed or outcrossed. Progeny are analyzed to identify those individuals having endogenous wild-type delta-12 fatty acid desaturase activity inhibited by dominant negative suppression as discussed above.

[0077] Methods similar to those described above are used to make delta-15 desaturase dominant negative suppression constructs, comprising a mutant delta-15 desaturase gene operably linked to at least one regulatory sequence. Transformation of a plant with such a construct will result in dominant negative suppression of endogenous delta-15 desaturase activity in transgenic progeny and in a decreased level of α -linolenic acid in homozygous dominant suppression lines. Such lines will have from about <1% to about 10% α -linolenic acid, preferably from about <1% to about 5%, based on total seed fatty acid composition.

[0078] In one embodiment of the invention, a plant contains a mutant delta-12 fatty acid desaturase and a mutant delta-15 fatty acid desaturase, both of which are expressed preferentially in seeds. Such a plant can be produced from the cross of single mutant plants, followed by outcrossing or selfing in order to obtain progeny seeds carrying both mutant chimeric genes. Progeny seeds are screened in order to identify those seeds carrying both mutant genes. Alternatively, seed-specific defects in delta-12 desaturase and delta-15 desaturase may be introduced into a wild-type plant by transformation, simultaneously or sequentially, with one or more dominant negative suppression constructs for delta-12 desaturase and delta-15 desaturase, each driven by suitable regulatory sequences. Other similar methods to construct double mutant plants will be recognized by those of skill in the art.

[0079] Double mutant plants can have more extreme fatty acid profiles in seeds than the single mutant plants, e.g., the double mutant phenotype can result in significantly lower levels of α -linolenic acid in seeds than the single mutant delta-15 desaturase plant phenotype. Thus, by combining seed-specific inhibition of microsomal delta-12 desaturase with seed-specific inhibition of microsomal delta-15 desaturase, one can obtain levels of seed α -linolenic acid that are as low or lower than those in a single mutant without adversely affecting desirable properties. The decreased levels of α -linolenic acid in the double homozygotes may be associated with increased levels of oleic acid and decreased levels of saturates and linoleic acid.

[0080] A preferred class of heterologous hosts for the expression of the nucleic acid fragments of the invention are eukaryotic hosts, particularly the cells of higher plants. Particularly preferred among the higher plants are the oil-producing species, such as soybean (Glycine max), rapeseed (including Brassica napus, B. campestris), sunflower (Helianthus annus), cotton (Gossypium hirsutum), corn (Zea mays), cocoa (Theobroma cacao), safflower (Carthamus tinctorius), oil palm (Elaeis guineensis), coconut palm (Cocos nucifera), flax (Linum usitatissimum), and peanut (Arachis hypogaea).

[0081] Expression in plants will use regulatory sequences functional in such plants. The expression of foreign genes in plants is well-established (De Blaere et al., *Meth. Enzymol.* (1987) 153:277-291). The source of the promoter chosen to drive the expression of the fragments of the invention is not critical provided it has sufficient transcriptional activity to accomplish the invention by increasing or decreasing, respectively, the level of translatable mRNA for the fatty acid desaturases in the desired host tissue. Preferred promoters include (a) strong constitutive plant promoters, such as those directing the 19S and 35S transcripts in cauliflower mosaic virus (Odell et al., *Nature* (1985) 313:810-812; Hull et al., *Virology* (1987) 86:482-493),(b) tissue- or developmentally-specific promoters, and (c) other transcriptional promoter systems engineered in plants, such as those using bacteriophage T7 RNA polymerase promoter sequences to express foreign genes. Examples of tissue-specific promoters are the light-inducible promoter of the small subunit of ribulose 1,5-bis-phosphate carboxylase (if expression is desired in photosynthetic tissues), the maize zein protein promoter (Matzke et al., *EMBO J.* (1984) 3:1525-1532), and the chlorophyll a/b binding protein promoter (Lampa et al., *Nature* (1986) 316:750-752).

[0082] Particularly preferred promoters are those that allow seed-specific expression. This may be especially useful since seeds are the primary source of vegetable oils and also since seed-specific expression will avoid any potential deleterious effect in non-seed tissues. Examples of seed-specific promoters include, but are not limited to, the promoters of seed storage proteins, which can represent up to 90% of total seed protein in many plants. The seed storage proteins are strictly regulated, being expressed almost exclusively in seeds in a highly tissue-specific and stage-specific manner (Higgins et al., *Ann. Rev. Plant Physiol.* (1984) 35:191-221; Goldberg et al., *Cell* (1989) 56:149-160). Moreover, different seed storage proteins may be expressed at different stages of seed development.

[0083] Expression of seed-specific genes has been studied in great detail (See reviews by Goldberg et al., Cell (1989) 56:149-160 and Higgins et al., Ann. Rev. Plant Physiol. (1984) 35:191-221). There are currently numerous examples of seed-specific expression of seed storage protein genes in transgenic dicotyledonous plants. These include genes from dicotyledonous plants for bean β-phaseolin (Sengupta-Gopalan et al., Proc. Natl. Acad. Sci. USA (1985) 82:3320-3324; Hoffman et al., Plant Mol. Biol. (1988) 11:717-729), bean lectin (Voelker et al., EMBO J. (1987) 6:3571-3577), soybean lectin (Okamuro et al., Proc. Natl. Acad. Sci. USA (1986) 83:8240-8244), soybean Kunitz trypsin inhibitor (Perez-Grau et al., Plant Cell (1989) 1:095-1109), soybean β-conglycinin (Beachy et al., EMBO J. (1985) 4:3047-3053; pea vicilin (Higgins et al., Plant Mol. Biol. (1988) 11:683-695), pea convicilin (Newbigin et al., Planta (1990) 180:461-470), pea legumin (Shirsat et al., Mol. Gen. Genetics (1989) 215:326-331); rapeseed napin (Radke et al., Theor. Appl. Genet. (1988) 75:685-694) as well as genes from monocotyledonous plants such as for maize 15 kD zein (Hoffman et al., EMBO J. (1987) 6:3213-3221), maize 18 kD oleosin (Lee et al., Proc. Natl. Acad. Sci. USA (1991) 888:6181-6185), barley β-hordein (Marris et al., Plant Mol. Biol. (1988) 10:359-366) and wheat glutenin (Colot et al., EMBO J. (1987) 6:3559-3564). Moreover, promoters of seed-specific genes operably linked to heterologous coding sequences in chimeric gene constructs also maintain their temporal and spatial expression pattern in transgenic plants. Such examples include use of Arabidopsis thaliana 2S seed storage protein gene promoter to express enkephalin peptides in Arabidopsis and B. napus seeds (Vandekerckhove et al., Biol Technology (1989) 7:929-932), bean lectin and bean β-phaseolin promoters to express luciferase (Riggs et al., Plant Sci. (1989) 63:47-57), and wheat glutenin promoters to express chloramphenicol acetyl transferase (Colot et al., EMBO J. (1987) 6:3559-3564).

[0084] Of particular use in the expression of the nucleic acid fragment of the invention will be the heterologous promoters from several soybean seed storage protein genes such as those for the Kunitz trypsin inhibitor (Jofuku et al., Plant Cell (1989) 1:1079-1093; glycinin (Nielson et al., Plant Cell (1989) 1:313-328), and β-conglycinin (Harada et al., Plant Cell (1989) 1:415-425). Promoters of genes for α- and β-subunits of soybean β-conglycinin storage protein will be particularly useful in expressing the mRNA or the antisense RNA in the cotyledons at mid- to late-stages of seed development (Beachy et al., EMBOJ. (1985) 4:3047-3053) in transgenic plants. This is because there is very little position effect on their expression in transgenic seeds, and the two promoters show different temporal regulation. The promoter for the α-subunit gene is expressed a few days before that for the β-subunit gene. This is important for transforming rapeseed where oil biosynthesis begins about a week before seed storage protein synthesis (Murphy et al., J. Plant Physiol. (1989) 135:63-69).

[0085] Also of particular use will be promoters of genes expressed during early embryogenesis and oil biosynthesis. The native regulatory sequences, including the native promoters, of the fatty acid desaturase genes expressing the nucleic acid fragments of the invention can be used following their isolation by those skilled in the art. Heterologous promoters from other genes involved in seed oil biosynthesis, such as those for <u>B. napus</u> isocitrate lyase and malate synthase (Comai et al., *Plant Cell* (1989) 1:293-300), delta-9 desaturase from safflower (Thompson et al. *Proc. Natl. Acad.*

Sci. USA (1991) 88:2578-2582) and castor (Shanklin et al., Proc. Natl. Acad. Sci. USA (1991) 88:2510-2514), acyl carrier protein (ACP) from Arabidopsis (Post-Beittenmiller et al., Nucl. Acids Res. (1989) 17:1777), B. napus (Safford et al., Eur. J. Biochem. (1988) 174:287-295), and B. campestris (Rose et al., Nucl. Acids Res. (1987) 15:7197), β-ketoacyl-ACP synthetase from barley (Siggaard-Andersen et al., Proc. Natl. Acad. Sci. USA (1991) 88:4114-4118), and oleosin from Zea mays (L e et al., Proc. Natl. Acad. Sci. USA (1991) 88:6181-6185), soybean (Genbank Accession No: X60773) and B. napus (Lee et al., Plant Physiol. (1991) 96:1395-1397) will be of use. If the sequence of the corresponding genes is not disclosed or their promoter region is not identified, one skilled in the art can use the published sequences for the relatively-abundant enoyl-ACP reductase and acetyl-CoA carboxylase are also published (Slabas et al., Biochim. Biophys. Acta (1987) 877:271-280; Cottingham et al., Biochim. Biophys. Acta (1988) 954:201-207) and one skilled in the art can use these sequences to isolate the corresponding seed genes with their promoters. Similarly, the fragments of the present invention encoding fatty acid desaturases can be used to obtain promoter regions of the corresponding genes for use in expressing chimeric genes.

[0086] Attaining the proper level of expression of the nucleic acid fragments of the invention may require the use of different chimeric genes utilizing different promoters. Such chimeric genes can be transferred into host plants either together in a single expression vector or sequentially using more than one vector.

[0087] It is envisioned that the introduction of enhancers or enhancer-like elements into the promoter regions of either the native or chimeric nucleic acid fragments of the invention will result in increased expression to accomplish the invention. This would include viral enhancers such as that found in the 355 promoter (Odell et al., *Plant Mol. Biol.* (1988) 10:263-272), enhancers from the opine genes (Fromm et al., *Plant Cell* (1989) 1:977-984), or enhancers from any other source that result in increased transcription when placed into a promoter operably linked to a nucleic acid fragment of the invention.

[0088] Of particular importance is the DNA sequence element isolated from the gene for the α -subunit of β -conglycinin that can confer 40-fold seed-specific enhancement to a constitutive promoter (Chen et al., *Dev. Genet.* (1989) 10:112-122). One skilled in the art can readily isolate this element and insert it within the promoter region of any gene in order to obtain seed-specific enhanced expression with the promoter in transgenic plants. Insertion of such an element in any seed-specific gene that is expressed at different times than the β -conglycinin gene will result in expression in transgenic plants for a longer period during seed development.

[0089] The invention can also be accomplished by a variety of other methods to obtain the desired end. In one form, the invention is based on modifying plants to produce increased levels of mutant fatty acid desaturases by virtue of introducing more than one copy of the foreign gene containing the nucleic acid fragments of the invention. In some cases, the desired level of polyunsaturated fatty acids may require introduction of foreign genes for more than one kind of mutant fatty acid desaturase.

[0090] Any 3' non-coding region capable of providing a polyadenylation signal and other regulatory sequences that may be required for the proper expression of the nucleic acid fragments of the invention can be used to accomplish the invention. This would include 3' ends of the native fatty acid desaturase(s), viral genes such as from the 35S or the 19S cauliflower mosaic virus transcripts, from the opine synthesis genes, ribulose 1,5-bisphosphate carboxylase, or chlorophyll a/b binding protein. There are numerous examples in the art that teach the usefulness of different 3' non-coding regions.

[0091] Various methods of transforming cells of higher plants according to the present invention are available to those skilled in the art (see EPO Pub. 0 295 959 A2 and 0 318 341 A1). Such methods include those based on transformation vectors utilizing the Ti and Ri plasmids of <u>Agrobacterium spp</u>. It is particularly preferred to use the binary type of these vectors. Ti-derived vectors transform a wide variety of higher plants, including monocotyledonous and dicotyledonous plants (Sukhapinda et al., *Plant Mol. Biol.* (1987) 8:209-216; Potrykus, *Mol. Gen. Genet.* (1985) 199:183). Other transformation methods are available to those skilled in the art, such as direct uptake of foreign DNA constructs (see EPO Pub. 0 295 959 A2), techniques of electroporation (Fromm et al., *Nature* (1986) (London) 319:791) or high-velocity ballistic bombardment with metal particles coated with the nucleic acid constructs (Kline et al., *Nature* (1987) (London) 327:70). Once transformed, the cells can be regenerated by those skilled in the art.

[0092] Of particular relevance are the recently described methods to transform foreign genes into commercially important crops, such as rapeseed (De Block et al., *Plant Physiol.* (1989) 91:694-701), sunflower (Everett et al., *Biol Technology* (1987) 5:1201), and soybean (Christou et al., *Proc. Natl. Acad. Sci* USA (1989) 86:7500-7504.

[0093] The use of restriction fragment length polymorphism (RFLP) markers in plant breeding has been well-documented in the art (Tanksley et al., Bio/Technology (1989) 7:257-264). Thus, the nucleic acid fragments of the invention can be used as molecular markers for traits associated with mutant fatty acid desaturases. These traits will include altered levels of unsaturated fatty acids. The nucleic acid fragment of the invention can also be used to isolate the fatty acid desaturase gene from other mutant plants with altered levels of unsaturated fatty acids. Sequencing of these genes will reveal nucleotide differences that cause the alteration in levels of unsaturated fatty acids. Oligonucleotides designed around these differences may also be used in plant breeding as diagnostic markers to follow fatty acid variation. In one

embodiment, oligonucleotides based on differences between wt and mutant $\Delta 12$ des may be used as molecular markers in breeding canola: lines with variant oil profiles.

EXAMPLES

[0094] The present invention is further defined in the following Examples, in which all parts and percentages are by weight and degrees are Celsius, unless otherwise stated. It should be understood that these Examples, while indicating preferred embodiments of the invention, are given by way of illustration only. From the above discussion and these Examples, one skilled in the art can ascertain the essential characteristics of this invention, and without departing from the spirit and scope thereof, can make various changes and modifications of the invention to adapt it to various usages and conditions. All publications, including patents and non-patent literature, referred to in this specification are expressly incorporated by reference herein.

EXAMPLE 1

SEQUENCES OF MUTANT DELTA-12 FATTY ACID DESATURASES FROM B. NAPUS

[0095] Primers specific for the FAD2 structural gene were used to clone the entire open reading frame (ORF) of the D and F forms of the gene by reverse transcription-polymerase chain reaction (RT-PCR). The sequences of the primers used for isolation of the D form ORF of B. napus FAD2 gene are

5'-CATGGGTGCAGGTGGAAGAATGC-3' (SEQ ID NO: 9); and 5'-GTTTCTTCTTTGCTTCATAAC-3' (SEQ ID NO: 10).

[0096] The sequences of the primers used to clone the F form ORF of B. napus FAD2 gene are

5'-CATGGGTGCAGGTGGAAGAATGC-3' (SEQ ID NO: 11); and 5'-TCTTTCACCATCATCATATCC-3' (SEQ ID NO: 12).

RNA from seeds of three lines, IMC129, Q508 and Westar, was isolated by an acid guanidinium thiocyanate-phenol-chloroform extraction method (Chomczynski and Sacchi, 1987; *Analytical Biochemistry* 162, 156-159, 1987). The total RNA was used as a template for reverse transcription and PCR amplification by RNA PCR kit (Perkin Elmer). The RT-PCR amplified fragments were cloned into pGEM-T vector (Promega), and then used for nucleotide sequence determination. The DNA sequence of each gene from each line was determined from both strands by dideoxy sequencing by Sanger et al. (Proc Natl Acad Sci USA 74, 5463-5467).

[0097] The D gene of IMC129 contained a G to A transversion at nucleotide 316 (from the translation initiation codon) of the D gene in IMC129 (SEQ ID NO:3), compared to the sequence of Westar. The transversion changes the codon at this position from GAG to AAG and results in a substitution of glutamic acid by lysine. The same base change was also detected in IMC129 when RNA from leaf tissue was used as template. The G to A mutation at nucleotide 316 was confirmed by sequencing several independent clones containing fragments amplified directly from genomic DNA of IMC129. These results eliminated the possibility of a rare mutation introduced during reverse transcription and PCR in the RT-PCR protocol. The mutation in the D form of delta-12 desaturase in IMC129 mapped to a conserved region of cloned delta-12 and delta-15 membrane bound-desaturases (Table 2).

[0098] The sequence of the F form of delta-12 desaturase in IMC129 was the same as the F form of delta-12 desaturase in Westar (SEQ ID NO:5).

[0099] For Q508, the sequence of the D form of delta-12 desaturase (SEQ ID NO:3) was the same as the D form of the IMC129 gene. This was expected, as Q508 was derived by mutagenesis of IMC129.

[0100] Sequence analysis of the Q508 F form of delta-12 desaturase revealed a T to A transition at nucleotide 515 (SEQ ID NO:7), compared to the wild-type Westar sequence. This mutation results in a change from a CTC codon to a CAC codon, substituting a histidine residue for the wild-type leucine residue.

TABLE 2

Alignment of Amino Acid Sequences of Cloned Canola Membrane Bound-Desaturases											
Desaturase Gene	Sequence ^a	Position ^b									
Canola-FAD2-D	H <u>E</u> C <u>G</u> H	110									
Canola-FAD2-F	H <u>E</u> CGH	110									
Canola FAD6 ^c	H <u>D</u> C <u>A</u> H	171									
Canola-FAD3 ^d	H <u>D</u> Ċ <u>G</u> H	97									
Canola-FAD7 ^e	H <u>D</u> CGH	126									

^aOne letter amino acid code; conservative substitutions are underlined

EXAMPLE 2

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25 GENE-SPECIFIC OLIGONUCLEOTIDE MARKERS FOR THE MUTANT AND WILD TYPE DELTA-12 FATTY ACID DESATURASE GENES

[0101] The D form of IMC129 fad2 gene contains a G to A transversion at nucleotide 316 from the translation initiation codon. Two short oligonucleotide upstream (5') primers, based on the single base change (G to A) between the D form of the IMC129 and wild type FAD2 genes, were designed. The sequences of the upstream (5') primers are as follows:

5' gene-specific primer for wild type FAD2-D:

5'-GTCTGGGTCATAGCCCACG-3'

(SEQ ID NO:13); and

5' gene-specific primer for IMC129 fad2-d:

5'-GTCTGGGTCATAGCCCACA-3'

(SEQ ID NO:14).

A common downstream (3') primer (SEQ ID NO: 10) specific for the D form of the FAD2 gene was used for both IMC129 and wild type FAD2 genes. These gene-specific primers were used in a DNA based PCR diagnostic assay to genotype plants carrying the mutant and/or wild type FAD2 genes.

- [0102] Genomic DNA was isolated from leaf tissue of IMC129 and Westar plants, and used as PCR templates. The PCR amplification assays were carried out by using a gene amplification kit (Perkin Elmer). The PCR conditions are as follows: denaturing temperature, 95°C for 1 min; annealing temperature, 52°C for 1 min; amplification temperature 72°C for 1 min. Total 20 PCR cycles were extended. Under these conditions primers SEQ ID NO:13 and SEQ ID NO:14 only amplified wild type FAD2-D and IMC 129 mutant fad2-d gene fragments, respectively.
 - [0103] The specificity of the gene-specific primers could be further improved by shortening the length of the primers and by replacing the base C with a T at the second position from the 3' end of the oligonucleotide PCR primer for FAD2-D (SEQ ID NO:13). The sequences of the modified upstream (5') oligonucleotide PCR primers are as follows:

5' modified gene-specific primer for wild type FAD2-D:

5'-CTGGGTCATAGCCCATG-3'

(SEQ ID NO:15); and

5' modified gene-specific primer for IMC129 fad2-d:

5'-CTGGGTCATAGCCCACA-3'

(SEQ ID NO:16).

The same common downstream (3') oligonucleotide primer (SEQ ID NO:10) was used for amplifying FAD2-D and fad2-55 d. with the modified primers, the genotype for FAD2-D and fad2-d could be consistently distinguished after extended 30 cycle of PCR amplification. Therefore, the DNA based PCR assay provided a simple and reliable method of genotypping B. Napus germplasms containing mutant and/or wild type FAD2 genes.

^bPosition in gene product of first amino acid

cFAD6 = Plastid delta-12

dFAD3 = Microsomal delta-15

⁶FAD7 = Plastid delta-15

EXAMPLE 3

CONSTRUCTS FOR DOMINANT NEGATIVE SUPPRESSION OF DELTA-12 FATTY ACID DESATURASE

[0104] The vector pZS212 was used to construct plasmids for dominant negative suppression experiments. One construct was prepared by inserting the full-length mutant D gene coding sequence (nucleotides 1 to 1155 of SEQ ID NO:3) in sense orientation between the phaseolin promoter and phaseolin 3' poly A region of plasmid pCW108. The pCW108 vector contains the bean phaseolin promoter and 3' untranslated region and was derived from the commercially available pUC18 plasmid (Gibco-BRL) via plasmids AS3 and pCW104. Plasmid AS3 contains 495 base pairs of the bean (Phaseolus vulgaris) phaseolin (7S seed storage protein) promoter starting with 5'-TGGTCTTTTGGT-3' followed by the entire 1175 base pairs of the 3' untranslated region of the same gene (see sequence descriptions in Doyle et al., (1986) J. Biol. Chem. 261:9228-9238 and Slightom et al., (1983) Proc. Natl. Acad. Sci. USA, 80:1897-1901. Further sequence description may be found in WO 9113993) cloned into the Hind III site of pUC18. The additional cloning sites of the pUC18 multiple cloning region (Eco RI, Sph I, Pst I and Sal I) were removed by digesting with Eco RI and Sal I, filling in the ends with Klenow and religating to yield the plasmid pCW104. A new multiple cloning site was created between the 495bp of the 5' phaseolin and the 1175bp of the 3' phaseolin by inserting a dimer of complementary synthetic oligonucleotides consisting of the coding sequence for a Nco I site (5'-CCATGG-3') followed by three filler bases (5'-TAG-3'), the coding sequence for a Sma I site (5'-CCCGGG-3'), the last three bases of a Kpn I site (5'-TAG-3'), a cytosine and the coding sequence for an Xba I site (5'-TCTAGA-3') to create the plasmid pCW108. This plasmid contains unique Nco I, Sma I, Kpn I and Xba I sites directly behind the phaseolin promoter.

[0105] The resulting 5'-phaseolin promoter-mutant fad2-phaseolin poly A-3' construct was excised and cloned between the EcoRI/Sall sites of pZS212, resulting in the plasmid designated pZPhMCFd2 (Figure 1). pZS212 is based on a vector which contains: (1) the chimeric gene nopaline synthase/neomycin phosphotransferase as a selectable marker for transformed plant cells (Brevan et al. (1984) *Nature* 304: 184-186), (2) the left and right borders of the T-DNA of the Ti plasmid (Brevan et al. (1984) *Nucl. Acids Res.* 12:8711-8720), (3) the E. coli lacZ α-complementing segment (Vieria and Messing (1982) *Gene* 19:259-267) with unique restriction endonuclease sites for Eco RI, Kpn I, Bam HI, and Sal I, (4) the bacterial replication origin from the <u>Pseudomonas</u> plasmid pVS1 (Itoh et al. (1984) *Plasmid* 11:206-220), and (5) the bacterial neomycin phosphotransferase gene from Tn5 (Berg et al. (1975) Proc. *Natnl. Acad. Sci.* U.S.A. 72:3628-3632) as a selectable marker for transformed <u>A. tumefaciens</u>. The nopaline synthase promoter in the plant selectable marker was replaced by the 35S promoter (Odell et al. (1985) *Nature*, 313:810-813) by a standard restriction endonuclease digestion and ligation strategy.

[0106] A second plasmid was constructed by inserting the full-length wild type canola D gene coding sequence (nucleotides 130 to 1281 of SEQ ID NO:1) into the Notl site of the canola napin promoter expression vector plMC401 which contains a 2.2 kb napin expression cassette.

[0107] The canola napin promoter expression cassette in pIMC401 was constructed as follows: ten oligonucleotide primers were synthesized based upon the nucleotide sequence of napin lambda clone CGN1-2 published in European Patent Application EP 255378). The oligonucleotide sequences were:

- BR42 and BR43 corresponding to bases 1132 to 1156 (BR42) and the complement of bases 2248 to 2271 (BR43) of the sequence listed in Figure 2 of EP 255378.
- BR45 and BR46 corresponding to bases 1150 to 1170 (BR46) and the complement of bases 2120 to 2155 (BR45) of the sequence listed in Figure 2 of EP 255378. In addition BR46 had bases corresponding to a Sal I site (5'-GTC-GAC-3') and a few additional bases (5'-TCAGGCCT-3') at its 5' end and BR45 had bases corresponding to a Bgl II site (5'-AGATCT-3') and two (5'-CT-3') additional bases at the 5' end of the primer,
- 45 BR47 and BR48 corresponding to bases 2705 to 2723 (BR47) and bases 2643 to 2666 (BR48) of the sequence listed in Figure 2 of EP 255378. In addition BR47 had two (5'-CT-3') additional bases at the 5' end of the primer followed by bases corresponding to a Bgl II site (5'-AGATCT-3') followed by a few additional bases (5'-TCAGGCCT-3').
 - BR49 and BR50 corresponding to the complement of bases 3877 to 3897 (BR49) and the complement of bases 3985 to 3919 (BR50) of the sequence listed in Figure 2 of EP 255378. In addition BR49 had bases corresponding to a Sal I site (5'-GTCGAC-3') and a few additional bases (5'-TCAGGCCT-3') at its 5' end,
 - BR57 and BR58 corresponding to the complement of bases 3875 to 3888 (BR57) and bases 2700 to 2714 (BR58) of the sequence listed in Figure 2 of EP 255378. In addition the 5' end of BR57 had some extra bases (5'-CCATGG-3') followed by bases corresponding to a Sac I site (5'-GAGCTC-3') followed by more additional bases (5'-GTC-GACGAGG-3'). The 5' end of BR58 had additional bases (5'-GAGCTC-3') followed by bases corresponding to a Nco I site (5'-CCATGG-3') followed by additional bases (5'-AGATCTGGTACC-3').
 - BR61 and BR62 corresponding to bases 1846 to 1865 (BR61) and bases 2094 to 2114 (BR62) of the sequence listed in Figure 2 of EP 255378. In addition the 5' end of BR 62 had additional bases (5'-GACA-3') followed by bases corresponding to a Bgl II site (5'-AGATCT-3') followed by a few additional bases (5'-GCGGCCGC-3').

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[0108] Genomic DNA from the canola variety 'Hyola401' (Zeneca Seeds) was used as a template for PCR amplification of the napin promoter and napin terminator regions. The promoter was first amplified using primers BR42 and BR43, and reamplified using primers BR45 and BR46. Plasmid plMC01 was derived by digestion of the 1.0 kb promoter PCR product with Sall/Bglll and ligation into Sall/BamHI digested pBluescript SK+ (Stratagene). The napin terminator region was amplified using primers BR48 and BR50, and reamplified using primers BR47 and BR49. Plasmid plMC06 was derived by digestion of the 1.2 kb terminator PCR product with Sall/Bglll and ligation into Sall/Bglll digested pSP72 (Promega). Using plMC06 as a template, the terminator region was reamplified by PCR using primer BR57 and primer BR58. Plasmid plMC101 containing both the napin promoter and terminator was generated by digestion of the PCR product with Sacl/Ncol and ligation into Sacl/Ncol digested plMC01. Plasmid plMC101 contains a 2.2 kb napin expression cassette including complete napin 5' and 3' non-translated sequences and an introduced Ncol site at the translation start ATG. Primer BR61 and primer BR62 were used to PCR amplify an. "270 bp fragment from the 3' end of the napin promoter. Plasmid plMC401 was obtained by digestion of the resultant PCR product with EcoRl/Bglll and ligation into EcoRl/Bglll digested plMC101. Plasmid plMC401 contains a 2.2 kb napin expression cassette lacking the napin 5' non-translated sequence and includes a Notl site at the transcription start.

[0109] The fragment containing the 5'-napin-fad2D-napin poly A-3' cassette was then inserted into the Sall site of pZS212, and the resulting 17.2 Kb plasmid was termed pIMC127 (Figure 2).

[0110] A third plasmid, plMC135, was constructed in a manner similar to that described above for plMC127. Plasmid plMC135 contains a 5' cruciferin promoter fragment operably linked in sense orientation to the full-length wild-type coding sequence of SEQ ID NO:1, followed by a cruciferin 3' poly A fragment.

[0111] A fourth plasmid, pIMC140 was constructed in a manner similar to that described above. Plasmid pIMC140 contains a 5' napin promoter fragment operably linked in sense orientation to the full-length mutant Q508 F gene coding sequence (SEQ ID NO:7), followed by a 3' napin poly A fragment.

EXAMPLE 4

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FATTY ACID PROFILES IN DOMINANT NEGATIVE SUPPRESSION PLANTS

[0112] The plasmids pZPhMCFd2, pIMC127, pIMC135 and pIMC140 were transferred by a freeze/thaw method (Holsters et al. (1978) *Mol Gen Genet* 163:181-187) to the <u>Agrobacterium</u> strain LBA4404/pAL4404 (Hockema et al. (1983), *Nature* 303:179-180).

[0113] Brassica <u>napus</u> cultivar "Westar" was transformed by co-cultivation of seedling pieces with disarmed <u>Agrobacterium tumefaciens</u> strain LBA4404 carrying the appropriate binary vector.

[0114] B. napus seeds were sterilized by stirring in 10% Chlorox, 0.1% SDS for thirty min, and then rinsed thoroughly with sterile distilled water. The seeds were germinated on sterile medium containing 30 mM CaCl₂ and 1.5% agar, and grown for six days in the dark at 24°C.

[0115] Liquid cultures of <u>Agrobacterium</u> for plant transformation were grown overnight at 28°C in Minimal A medium containing 100 mg/L kanamycin.

Minimal A Bacterial Growth Medium

[0116] Dissolve in distilled water:

10.5 grams potassium phosphate, dibasic
4.5 grams potassium phosphate, monobasic
1.0 gram ammonium sulfate
0.5 gram sodium citrate, dihydrate
Make up to 979 mL with distilled water
Autoclave
Add 20 mL filter-sterilized 10% sucrose
Add 1 mL filter-sterilized 1 M MgSO₄

[0117] The bacterial cells were pelleted by centrifugation and resuspended at a concentration of 10^8 cells/mL in liquid Murashige and Skoog Minimal Organic medium (GIBCO; Cat. No. 510-3118) containing 100 μ M acetosyringone. [0118] B. napus seedling hypocotyls were cut into 5 mm segments which were immediately placed into the bacterial suspension. After 30 min, the hypocotyl pieces were removed from the bacterial suspension and placed onto BC-35 callus medium containing 100 μ M acetosyringone.

Brassica Callus Medium BC-35

[0119] Per liter:

5 Murashige and Skoog Minimal Organic Medium (MS salts, 100 mg/L i-inositol, 0.4 mg/L thiamine;

GIBCO #510-3118)

30 grams sucrose

18 grams mannitol

0.5 mg/L 2,4-D

0.3 mg/L kinetin

0.6% agarose

pH 5.8

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The plant tissue and Agrobacteria were co-cultivated for three days at 24°C in dim light.

[0120] The co-cultivation was terminated by transferring the hypocotyl pieces to BC-35 callus medium containing 200 mg/L carbenicillin to kill the <u>Agrobacteria</u>, and 25 mg/L kanamycin to select for transformed plant cell growth. The seed-ling pieces were incubated on this medium for three weeks at 28°C under continuous light.

[0121] After four weeks, the segments were transferred to BS-48 regeneration medium containing 200 mg/L carbenicillin and 25 mg/L kanamycin.

Brassica Regeneration Medium BS-48

[0122]

25 Murashige and Skoog Minimal Organic Medium

Gamborg B5 Vitamins (SIGMA #1019)

10 grams glucose

250 mg xylose

600 mg MES

0.4% agarose

pH 5.7

Filter-sterilize and add after autoclaving:

2.0 mg/L zeatin

0.1 mg/L IAA

[0123] Plant tissue was subcultured every two weeks onto fresh selective regeneration medium, under the same culture conditions described for the callus medium. Putatively transformed calli grew rapidly on regeneration medium; as calli reached a diameter of about 2 mm; they were removed from the hypocotyl pieces and placed on the same medium lacking kanamycin.

[0124] Shoots began to appear within several weeks after transfer to BS-48 regeneration medium. As soon as the shoots formed discernable stems, they were excised from the calli, transferred to MSV-1A elongation medium, and moved to a 16:8 h photoperiod at 24°C.

45 Brassica Shoot Elongation Medium MSV-1A

[0125]

Murashige and Skoog Minimal Organic Medium Gamborg B5 Vitamins 10 grams sucrose 0.6% agarose pH 5.8

55 [0126] Once shoots had elongated several internodes, they were cut above the agar surface and the cut ends were dipped in Rootone. Treated shoots were planted directly into wet Metro-Mix 350 soiless potting medium. The pots were covered with plastic bags which were removed when the plants were clearly growing -- after about ten days.

[0127] Plants were grown under a 16:8 h photoperiod, with a daytime temperature of 23°C and a nightime tempera-

ture of 17°C. When the primary flowering stem began to elongate, it was covered with a mesh pollen-containment bag to prevent outcrossing. Self-pollination was facilitated by shaking the plants several times each day.

[0128] Transgenic progeny plants containing pZPhMCFd2 were designated as the WS201 series. Plants transformed with pIMC127 were designated as the WS127 series. Plants transformed with pIMC135 were designated as the WS135 series. Plants transformed with pIMC140 were designated as the WS140 series. Seeds were obtained by selfing the T1 plants. Fatty acid profiles of the T2 seeds were determined as described in WO 93/11245. The results are shown in Tables 3 and 4 and Figures 3 and 4.

T2 Se	T2 Seeds having Decreased C18:2 (WS201 ¹ Transformants) ² .														
, , , , , , , , , , , , , , , , , , ,			Fatty Ac	id Compo	sition(%)										
Line No.	Plant Series	C16:0	C18:0	C18:1	C18:2	C18:3									
T300236	WS201	4.1	3.2	72.5	10.6	6.5									
T300238	WS201	3.9	2.7	74.4	9,2	6.6									
T300390	WS201	4.2	2.5	75.4	9.3	5.4									
T300273	WS201	3.9	2.8	77.4	7.5	5.3									
T300400	WS201	3.9	2.5	78.6	6.6	5.3									
T300354	WS201	4.0	2.7	78.6	6.4	4.51									

¹ WS201: phaseolin promoter/IMC129 mutant, sense orientation.

[0129] As shown in Figure 3, a large proportion of the control WS127 plants have elevated levels of linoleic acid. This result is expected, since the delta-12 desaturase gene dosage is higher due to the extra copy of the wild-type delta-12 desaturase gene. A small proportion of WS127 plants show lower levels of linoleic acid, due to cosuppression.

[0130] In contrast, no WS201 plants had elevated levels of linoleic acid. This result confirms that the mutant delta-12 desaturase D form gene in WS201 plants is non-functional. Furthermore, a higher proportion of WS201 plants have decreased C18:2 (5-14%) compared to the proportion of WS127 control plants having decreased C18:2. The proportion of WS201 plants having decreased C18:2 is about 25-fold higher than the proportion of WS127 plants having decreased C18:2. The surprising inhibition of endogenous delta-12 desaturase activity by a mutant delta-12 gene product has been termed dominant negative suppression.

[0131] The fatty acid composition of seeds produced by representative dominant negative suppression T2 WS201 plants is shown in Table 3. The plants show an altered fatty acid composition, including decreased C18:2, increased C18:1, and decreased saturates.

[0132] A T2 generation plant is not homozygous for the introduced gene. Consequently, T3 and subsequent generations that are homozygous for the mutant delta-12 desaturase gene will have even lower levels of linoleic acid, from about 1% to about 10%, preferably from about 1% to about 6%. Levels of oleic acid in homozygous lines will be from 75% to about 88%, preferably from about 80% to about 88%.

[0133] The results observed with WS140 plants (containing a mutant Q508 F form delta-12 desaturase gene) are shown in Figure 4. None of the WS140 plants have elevated C18:2 levels, similar to the results obtained with WS201 transgenic plants. As expected, a large proportion of the control WS135 plants have elevated C18:2 levels. The proportion of WS140 and WS135 plants having decreased C18:2 levels is similar, indicating that expression of this particular mutant delta-12 desaturase gene product does not inhibit endogenous wild-type delta-12 desaturase gene product to an extent greater than that expected from cosuppression. The fatty acid composition of seeds produced by a representative T2 WS140 plant with decreased C18:2 levels is shown in Table 4.

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² Population of 168 selfed individuals from greenhouse.

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TABLE 4

T2 Seed Having Decreased C18:2 (WS140 ¹ Transformants) ² .															
			Fatty Acid Composition (%)												
Line No.	Vector	C16:0	C18:0	C18:1	C18:2	C18:3									
T300435	pIMC140	3.8	3.5	73.7	9.7	6.11									

^{1.}WS140: napin promoter/Q508 fad2 F mutation, sense orientation

² Population of 61 selfed individuals from greenhouse

SEQUENCE LISTING

5	(1) GENERAL INFORMATION:	
10	 (i) APPLICANT: (A) NAME: E.I. Du Pont De Nemours and Company (B) STREET: 1007 Market Street (C) CITY: Wilmington (D) STATE: Delaware (E) COUNTRY: USA 	
	(F) POSTAL CODE (ZIP): 19898 (ii) TITLE OF INVENTION: Process	
15	(iii) NUMBER OF SEQUENCES: 16	
20	 (iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: Patentin Release #1.0, Version #1.30 (EPO) 	
25	(2) INFORMATION FOR SEQ ID NO:1: (i) SEQUENCE CHARACTERISTICS:	
30	(A) LENGTH: 1464 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA	
35	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1301281	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
40	GGCACGAGCT CGTGCCGAAT TCGGCACGAG AGGAGACAGA GAGAGAGTTT GAGGAGGAGC TTCTTCGTAG GGTTCATCGT TATTAACGTT AAATCTTCAT CCCCCCCTAC GTCAGCCAGC	120
45	TCAAGAAAC ATG GGT GCA GGT GGA AGA ATG CAA GTG TCT CCT CCC TCC Met Gly Ala Gly Gly Arg Met Gln Val Ser Pro Pro Ser 1 5 10	168
	AAA AAG TCT GAA ACC GAC AAC ATC AAG CGC GTA CCC TGC GAG ACA CCG Lys Lys Ser Glu Thr Asp Asn Ile Lys Arg Val Pro Cys Glu Thr Pro 15 20 25	216
50	CCC TTC ACT GTC GGA GAA CTC AAG AAA GCA ATC CCA CCG CAC TGT TTC Pro Phe Thr Val Gly Glu Leu Lys Lys Ala Ile Pro Pro His Cys Phe 30 35 40 45	264

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5					TCT Ser							312
	_	_			TAC Tyr							360
10	CCT Pro				TTC Phe							408
15					GTC Val 100						 	456
20					CAG Gln							504
20					GTC Val							552
<i>25</i>	CGA Arg											600
30	GTC Val									Tyr		648
35	AAC Asn							_				696
	TGG Trp 190				TTC Phe							744
40	GGC Gly				CAC His							792
4 5					ATC Ile							840
			Tyr		GCT Ala				_			888
50					CTT Leu 260							936

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	ATC ACT TAC TTG CAG CAC ACG CAT CCT TCC CTG CCT CAC TAT GAC TCG Ile Thr Tyr Leu Gln His Thr His Pro Ser Leu Pro His Tyr Asp Ser 270 285	984												
5	TCT GAG TGG GAT TGG TTG AGG GGA GCT TTG GCC ACC GTT GAC AGA GAC Ser Glu Trp Asp Trp Leu Arg Gly Ala Leu Ala Thr Val Asp Arg Asp 290 295 300	.032												
10	TAC GGA ATC TTG AAC AAG GTC TTC CAC AAT ATC ACG GAC ACG CAC GTG Tyr Gly Ile Leu Asn Lys Val Phe His Asn Ile Thr Asp Thr His Val 305 310 315	.080												
15	GCG CAT CAC CTG TTC TCG ACC ATG CCG CAT TAT CAT GCG ATG GAA GCT Ala His His Leu Phe Ser Thr Met Pro His Tyr His Ala Met Glu Ala 320 325 330	.128												
	ACG AAG GCG ATA AAG CCG ATA CTG GGA GAG TAT TAT CAG TTC GAT GGG Thr Lys Ala Ile Lys Pro Ile Leu Gly Glu Tyr Tyr Gln Phe Asp Gly 335 340 345	.176												
20	ACG CCG GTG GTT AAG GCG ATG TGG AGG GAG GCG AAG GAG TGT ATC TAT Thr Pro Val Val Lys Ala Met Trp Arg Glu Ala Lys Glu Cys Ile Tyr 350 365	224												
25	GTG GAA CCG GAC AGG CAA GGT GAG AAG GGT GTG TTC TGG TAC AAC Val Glu Pro Asp Arg Gln Gly Glu Lys Lys Gly Val Phe Trp Tyr Asn 370 375 380	272												
	AAT AAG TTA TGAAGCAAAG AAGAAACTGA ACCTTTCTCT TCTATGATTG Asn Lys Leu													
30	TCTTTGTTTA AGAAGCTATG TTTCTGTTTC AATAATCTTA ATTATCCATT TTGTTGTGTT 1													
	TTCTGACATT TTGGCTAAAA TTATGTGATG TTGGAAGTTA GTGTCTAAAA AAAAAAAAAA	1464												
35	(2) INFORMATION FOR SEQ ID NO:2:													
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 384 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear													
	(ii) MOLECULE TYPE: protein													
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:													
	Met Gly Ala Gly Gly Arg Met Gln Val Ser Pro Pro Ser Lys Lys Ser 1 1 15													
50	Glu Thr Asp Asn Ile Lys Arg Val Pro Cys Glu Thr Pro Pro Phe Thr 20 25 30													
	Val Gly Glu Leu Lys Lys Ala Ile Pro Pro His Cys Phe Lys Arg Ser 35 40 45													

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	Ile	Pro 50	Arg	Ser	Phe	Ser	Tyr 55	Leu	Ile	Trp	Asp	Ile 60	Ile	Ile	Ala	Ser
5	Сув 65	Phe	Tyr	Tyr		Ala 70	Thr	Thr	Tyr	Phe	Pro 75	Leu	Leu	Pro	His	Pro 80
	Leu	Ser	Tyr	Phe-	Ala 85	Trp.	Pro 	Leu	Tyr	Trp 90	Ala	Сув	Gln	Gly	Сув 95	Val
10	Leu	Thr	Gly	Val 100	Trp		Ile		His 105	Glu	Сув	Glу	His	His 110	Ala	Phe
15	Ser	qaA	Tyr 115	Gln	Trp	Leu	qaA	Asp 120	Thr	Val	Gly	Leu	Ile 125	Phe	His	Ser
	Phe	Leu 130	Leu	Val	Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140	His	Arg	Arg	His
	His 145	Ser	Asn	Thr	Gly	Ser 150	Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Lys 160
	Lys	Lys	Ser	Asp	Ile 165	Lys	Trp	Tyr	Gly	Lys 170	Tyr	Leu	Asn	Asn	Pro 175	Leu
25	Gly	Arg	Thr	Val 180	Met	Leu	Thr	Val	Gln 185	Phe	Thr	Leu	Gly	Trp 190	Pro	Leu
	Tyr		Ala 195		Asn	Val		Gly 200	Arg	Pro	Tyr	A ap	Gly 205	Gly	Phe	Ala
30	Сув	His 210	Phe	His	Pro	Asn			Ile	Tyr	Asn	Asp 220	Arg	Glu	Arg	Leu
	Gln 225	Ile	Tyr	Ile	Ser	А вр 230	Ala	Gly	Ile	Leu	Ala 235	Val	Суз	Tyr	Gly	Leu 240
35	Tyr	Arg	Tyr	Ala	Ala 245	Val	Gln	Gly	Val	Ala 250	Ser	Met	Val	Сув	Phe 255	Tyr
	Gly	Val	Pro	Leu 260	Leu	Ile	Val	Asn	Gly 265	Phe	Leu	Val	Leu	Ile 270	Thr	Tyr
40	Leu	Gln	His 275		His	Pro	Ser	Leu 280	Pro	His	Tyr	Asp	Ser 285	Ser	Glu	Trp
4 5	Asp	Trp 290	Leu	Arg	Gly	Ala	Leu 295	Ala	Thr	Val	qaA	Arg 300	qaA	Tyr	Gly	Ile
	Leu 305	Asn	Lys	Val	Phe	His 310	Asn	Ile	Thr	Asp	Thr 315	His	Val	Ala	His	His 320
50	Leu	Phe	Ser	Thr	Met 325	Pro	His	Tyr	His	Ala 330	Met	Glu	Ala	Thr	Lys 335	Ala
	Ile	Lys	Pro	Ile 340	Leu	Gly	Glu	Tyr	Tyr 345	Gln	Phe	Asp	Gly	Thr 350	Pro	Val

	Val Lys Ala Met Trp Arg Glu Ala Lys Glu Cys Ile Tyr Val 355 360 365	Glu Pro
5	Asp Arg Gln Gly Glu Lys Lys Gly Val Phe Trp Tyr Asn Asn 370 375 380	Lys Leu
	(2) INFORMATION FOR SEQ ID NO:3:	
10	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1155 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
15	(ii) MOLECULE TYPE: DNA	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
20	(vi) ORIGINAL SOURCE: (A) ORGANISM: Brassica napus	-
	(vii) IMMEDIATE SOURCE: (B) CLONE: IMC129	
25	(ix) FEATURE:	
	(D) OTHER INFORMATION: G to A transversion mutation at nucleotide 3 of the D form	16
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	**
	ATG GGT GCA GGT GGA AGA ATG CAA GTG TCT CCT CCC TCC AAA Met Gly Ala Gly Gly Arg Met Gln Val Ser Pro Pro Ser Lys	
	1 5 10	15
35	GAA ACC GAC AAC ATC AAG CGC GTA CCC TGC GAG ACA CCG CCC	
	Glu Thr Asp Asn Ile Lys Arg Val Pro Cys Glu Thr Pro Pro 20 25 30	Pne Thr
40	GTC GGA GAA CTC AAG AAA GCA ATC CCA CCG CAC TGT TTC AAA	
	Val Gly Glu Leu Lys Lys Ala Ile Pro Pro His Cys Phe Lys 35 40 45	arg ser
	ATC CCT CGC TCT TTC TCC TAC CTC ATC TGG GAC ATC ATA	
45	Ile Pro Arg Ser Phe Ser Tyr Leu Ile Trp Asp Ile Ile 50 55 60	Ala Ser
	TGC TTC TAC TAC GTC GCC ACC ACT TAC TTC CCT CTC CTC	CAC CCT 240
	Cys Phe Tyr Tyr Val Ala Thr Thr Tyr Phe Pro Leu Leu Pro 65 70 75	
50	CTC TCC TAC TTC GCC TGG CCT CTC TAC TGG GCC TGC CAG GGC	TGC GTC 288
	Leu Ser Tyr Phe Ala Trp Pro Leu Tyr Trp Ala Cys Gln Gly 85 90	

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O

5					TGG Trp													336
					TGG Trp												··.	384
10	-				CCT Pro													432
15					GGC Gly												•	480
20					ATC Ile 165												•	528
					ATG Met													576 ·
25	Tyr	Leu	Ala 195	Phe	AAC Asn	Val	Ser	Gly 200	Arg	Pro	Tyr	Asp	Gly 205	Gly	Phe	Ala		624
30	Сув	His 210	Phe	His	CCC Pro	Asn	Ala 215	Pro	Ile	Tyr	Asn	Asp 220	Arg	Glu	Arg	Leu 		672
35	Gln 225	Ile	Tyr	Ile	TCC Ser	Авр 230	Ala	Gly	Ile	Leu	Ala 235	Val	Сув	Tyr	Gly	Leu 240		720
	Tyr	Arg	Tyr	Ala	GCT Ala 245	Val	Gln	Gly	Val	Ala 250	Ser	Met	Val	Сув	Phe 255	Tyr		768
40	Gly	Val	Pro	Leu 260	CTG Leu	Ile	Val	Asn	Gly 265	Phe	Leu	Val	Leu	11e 270	Thr	Tyr		816
45	Leu	Gln	His 275	Thr	CAT His	Pro	Ser	Leu 280	Pro	His	Tyr	Asp	Ser 285	Ser	Glu	Trp		864
50	Aap	Trp 290	Leu	Arg	Gly	Ala	Leu 295	Ala	Thr	Val	Asp	Arg 300	Asp	Tyr	Gly	Ile		912
							Asn									CAC His 320		960

24

	CTG TTC TCG ACC ATG CCG CAT TAT CAT GCG ATG GAA GCT ACG AAG GCG Leu Phe Ser Thr Met Pro His Tyr His Ala Met Glu Ala Thr Lys Ala 325 330 335
5	ATA AAG CCG ATA CTG GGA GAG TAT TAT CAG TTC GAT GGG ACG CCG GTG 1056 11e Lys Pro 11e Leu Gly Glu Tyr Tyr Gln Phe Asp Gly Thr Pro Val 340 345 350
10	GTT AAG GCG ATG TGG AGG GAG GCG AAG GAG TGT ATC TAT GTG GAA CCG Val Lys Ala Met Trp Arg Glu Ala Lys Glu Cys Ile Tyr Val Glu Pro 355 360 365
15	GAC AGG CAA GGT GAG AAG AAA GGT GTG TTC TGG TAC AAC AAT AAG TTA T 1153 Asp Arg Gln Gly Glu Lys Lys Gly Val Phe Trp Tyr Asn Asn Lys Leu 370 375 380
	GA 1155
20	(2) INFORMATION FOR SEQ ID NO:4:
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 384 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear
25	(ii) MOLECULE TYPE: protein
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
30	Met Gly Ala Gly Gly Arg Met Gln Val Ser Pro Pro Ser Lys Lys Ser 1 10 15
	Glu Thr Asp Asn Ile Lys Arg Val Pro Cys Glu Thr Pro Pro Phe Thr 20 25 30
35	Val Gly Glu Leu Lys Lys Ala Ile Pro Pro His Cys Phe Lys Arg Ser 35 40 45
	Ile Pro Arg Ser Phe Ser Tyr Leu Ile Trp Asp Ile Ile Ile Ala Ser 50 55 60
40	Cys Phe Tyr Tyr Val Ala Thr Thr Tyr Phe Pro Leu Leu Pro His Pro 65 70 75 80
	Leu Ser Tyr Phe Ala Trp Pro Leu Tyr Trp Ala Cys Gln Gly Cys Val 85 90 95
45	Leu Thr Gly Val Trp Val Ile Ala His Lys Cys Gly His His Ala Phe 100 105 110
50	Ser Asp Tyr Gln Trp Leu Asp Asp Thr Val Gly Leu Ile Phe His Ser 115 120 125
	Phe Leu Leu Val Pro Tyr Phe Ser Trp Lys Tyr Ser His Arg Arg His 130 135 140

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	His 145	Ser	Asn	Thr	Gly	Ser 150	Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Lys 160
5	Lys	Lys	Ser	Ąsp	Ile 165	Lys	Trp	Tyr	Gly	Lys 170	Tyr	Leu	Asn	Asn	Pro 175	Leu
	Gly	Arg		180	Met		Thr	Val	Gln 185	Phe	Thr	Leu	Gly	Trp 190	Pro	Leu
10	Tyr	Leu	Ala 195	Phe	Asn	Val	Ser .,	-	Arg	Pro	Tyr	Asp	Gly 205	Gly	Phe	Ala
15	Сув	His 210	Phe	His	Pro	Asn	Ala 215	Pro	Ile	Tyr	Asn	Asp 220	Arg	Glu	Arg	Leu
	Gln 225	Ile	Tyr	Ile	Ser	Asp 230	Ala	Gly	Ile	Leu	Ala 235	Val	Сув	Tyr	Gly	Leu 240
20	Tyr	Arg	Tyr	Ala	Ala 245		Gln	Gly	Val	Ala 250	Ser	Met	Val	Сув	Phe 255	Tyr
	Gly	Val	Pro	Leu 260	Leu	Ile	Val	Asn	Gly 265	Phe	Leu	Val	Leu	Ile 270	Thr	Tyr
25	Leu	Gln	His 275	Thr	His	Pro	Ser	Leu 280	Pro	His	Tyr	Asp	Ser 285	Ser	Glu	Trp
	Asp	Trp 290	Leu	Arg	Gly	Ala	Leu 295	Ala	Thr	Val	Asp	Arg 300	Asp	Tyr	Gly	Ile
30	Leu 305	Asn	ГÀв	Val	Phe	His 310	Asn	Ile	Thr	Asp	Thr 315	His	Val	Ala	His	His 320
	Leu	Phe	Ser	Thr	Met 325	Pro	His	Tyr	His	Ala 330	Met	Glu	Ala	Thr	Lys 335	Ala
35	Ile	Lys	Pro	11e 340	Leu	Gly	Glu	Tyr	Tyr 345	Gln	Phe	qeA	Gly	Thr 350	Pro	Val
	Val	Lys	Ala 355	Met	Trp	Arg	Glu	Ala 360	Lys	Glu	Сув	Ile	Tyr 365	Val	Glu	Pro
40	Asp	Arg 370	Gln	Gly	Glu	Lys	Lys 375	Gly	Val	Phe	Trp	Tyr 380	Asn	Asn	Lys	Leu
45	(2)	I	NFORI	MATIC	ON FO	OR SI	EQ II	D NO	:5:							
	(:	•			CHAI				: pair:	3						
50			(B)	TYPI STR	E: 1 ANDEI OLOGY	nucle ONES	eic a	acid sing:								
	(i:	i.) 1	MOLE	CULE	TYP	E:	DNA							•		

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	(iii)	HYPOTH	HETICAL	: NO	>										
5	(iv)	ANTI-S	SENSE :	NO											
	(vi)		NAL SOU												
		(A) O	RGANISI	1: B	rass	ica	napu	s							
	(ix)	FEATUR	RE: OTHER II	JEÓÐM	IATTO	n.::	wild	Ltvn	e F	form	١.				
10											••				
	(xi)	_	NCE DES			_	_	NO:							
	ATG GGT Met Gly														48
15	net Gly	AIA G	5 SIY GIY	MY	Met	GIII	vai	10	210	210	JCI		15	5 C 2	
	GAA ACC														96
	Glu Thr	Asp T	Thr Ile 20	Lys	Arg	Val	Pro 25	Сув	Glu	Thr	Pro	Pro 30	Phe	Thr	
20	GTC GGA		ማር አእር	222	CCN	እ ምር	CCV	CCG	CAC	тст	TTC	ΔΔΔ	CGC	TCG	144
	Val Gly														
		35				40					45				
25	ATC CCT														192
	Ile Pro		ser Pne	ser	55	ьец	116	IID	Авр	60	116	116	AIG	Ser	
	TGC TTC	TAC T	TAC GTC	GCC	ACC	ACT	TAC	TTC	CCT	CTC	CTC	CCT	CAC	CCT	240
30	Cys Phe	Tyr T	Tyr Val	Ala 70	Thr	Thr	Tyr	Phe	Pro 75	Leu	Leu	Pro	His	Pro 80	
	65														
	CTC TCC														288
		-7-	85				•	90		-		_	95		
35	CTA ACC														336
	Leu Thr	_	Jal Trp 100	Val	Ile	Ala	His 105	Glu	Cys	Gly	His	His 110	Ala	Phe	
										oma.	3.000	mm.c	C) C	TOC .	384
40	AGC GAC														304
		115				120					125				
	TTC CTC														432
45	Phe Leu 130		Val Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140	Hls	Arg	ser	HIS	
	CAT TCC		act ccc	ጥሮሮ	CTC	GAG	AGA	GAC	GAA	GTG	Lalab	GTC	ccc	AAG	480
	His Ser								Glu					Lys	
50	145			150					155					160	
50	AAG AAG														528
	Lys Lys	s ser A	165		ιτb	ıyr	GIY	170	171	Leu.	TOIL		175		

5			ATG Met			_			_				576
			AAC Asn	_					_				624
10			CCC Pro										672
15			TCC Ser			_							720
20			GCC Ala 245										768
			CTG Leu										816
25			CAT His							_			864
30			GGA Gly				_						912
35			TTC Phe										960
40			ATG Met 325										1008
·			CTG Leu		Glu	Tyr	Gln						1056
45			TGG Trp										1104
50			GAG Glu								TTA Leu	T	1153
	GA												1155

		(2)	· II	NFOR	OITAN	ON FO	OR SI	II QE	NO:	6:							
5	Ir T	i)		SEQUI (A) (B) (D)	TYPE	: HT	384 mino	ami	no a		3						
10		(ii	.). 1	MOLE	CULE	TYPI	B: 1	prote	ein								
10		(xi	.) :	SEQUI	ENCE	DESC	CRIP	rion	: SI	EQ II	ON C	:6: ,.	• •		•		
		Met 1	Gly	Ala	Gly	Gly 5	Arg	Met	Gln	Val	Ser 10	Pro	Pro	Ser	Lys	Lys 15	Ser
15		Glu	Thr	Ąsp	Thr 20	Ile	Lys	Arg	Val	Pro 25	Cys	Glu	Thr	Pro	Pro 30	Phe	Thr
20		Val	Gly	Glu 35	Leu	ГÀВ	Lys	Ala	Ile 40	Pro	Pro	His	Сув	Phe 45	Lys	Arg	Ser
		Ile	Pro 50	Arg	Ser	Phe	Ser	Tyr 55	Leu	Ile	Trp	qaA	Ile 60	Ile	Ile	Ala	Ser
25		Сув 65	Phe	Tyr	Tyr	Val	Ala 70	Thr	Thr	Tyr	Phe	Pro 75	Leu	Leu	Pro	His	Pro 80
		Leu	Ser	Tyr	Phe	Ala 85	Trp	Pro	Leu	Tyr	Trp 90	Ala	Сув	Gln	Gly	Cys 95	Val
30		Leu	Thr	Gly	Val 100	Trp	Val	Ile	Ala	His 105	Glu	Сув	Gly	His	His 110	Ala	Phe
		Ser	Asp	Tyr 115	Gln	Trp	Leu	Asp	Asp 120	Thr	Val	Gly	Leu	Ile 125	Phe	His	Ser
35		Phe	Leu 130	Leu	Val	Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140	His	Arg	Ser	His
		His 145	Ser	Asn	Thr	Gly	Ser 150	Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Lys 160
40		Lys	Lys	Ser	qaA	Ile 165	Lys	Trp	Tyr	Gly	Lys 170	Tyr	Leu	Asn	Asn	Pro 175	Leu
45		Gly	Arg	Thr	Val 180	Met	Leu	Thr	Val	Gln 185	Phe	Thr	Leu	Gly	Trp 190	Pro	Leu
		Tyr	Leu	Ala 195	Phe	Asn	Va1	Ser	Gly 200	Arg	Pro	Tyr	Asp	G1y 205	Gly	Phe	Arg
50		Cys	His 210		His	Pro	Asn	Ala 215		Ile	Tyr	Asn	Asp 220	Arg	Glu	Arg	Leu
		Gln 225	Ile	Tyr	Ile	Ser	Asp 230		Gly	Ile	Leu	Ala 235		Сув	Tyr	Gly	Leu 240

	Phe	Arg	Tyr	Ala	Ala 245	Gly	Gln	Gly	Val	Ala 250	Ser	Net	Val	Сув	Phe 255	Tyr
5	Gly	Val	Pro	Leu. 260	Leu	Ile	Val	Asn _.	Gly 265	Phe	Leu	Val	Leu	Ile 270	Thr	Tyr
10	Leu	Gln	His 275					Leu 280		His	Tyr	Asp	Ser 285		Glu	_
	Asp	Trp 290	Phe	Arg	Gly	Ala	Leu 295	Ala	Thr	Val	Asp	Arg 300	Asp	Tyr	Ğly	Ile
15	Leu 305	As'n	Ļys	Val	Phe	His 310	Asn	Ile	Thr	Asp	Thr 315	His	Val	Ala	His	His 320
	Pro	Phe	Ser	Thr	Met 325	Pro	His	Tyr	His	Ala 330	Met	Glu	Ala	Thr	Lys 335	Ala
20	Ile	Lys	Pro	11e 340	Leu	Gly	Glu	Tyr	Tyr 345	Gln	Phe	Ąsp	Gly	Thr 350	Pro	Val
05	Val	Lys	Ala 355	Met	Trp	Arg		Ala .360	Lys	Glu	Сув	Ile	Tyr 365	Val	Glu	Pro
<i>25</i>	Asp	Arg 370	Gln	Gly	Glu	Lys	Lys 375	Gly	Val	Phe	Trp	Tyr 380	Asn	Asn	Lys	Leu
30	(2)	n	NFOR	(ATIC	ON FO	OR S	EQ II	ONO:	:7:							1
35	(i	ŕ	SEQUI (A) (B) (C) (D)	LENG TYPE STRA	TH:	11! nucle ONES!	55 ba	ingl	airs	3						
	(ii	L) 1	MOLE	CULE	TYP	E: 1	DNA									
40	(iii		HYPO' ANTI			: N	0									
45	(vi	•	ORIG (A)				Brası	sica	napı	18						
	(vii	L)	IMME (B)					В								
50	(i)	c)	FEAT		ER II	NFOR	MATIO	ON:	muta	atio		nuc		ide !	515	

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(xi)	SEQUENCE	DESCRIPTION:	SEO	ID :	NO:7:	
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5		GGT Gly															4	8
		ACC Thr					_										9	6
10	•==			20			_							: 30			• • • • • • • • • • • • • • • • • • • •	
	GTC																14	4
	Val	Gly	Glu 35	Leu	•	-		40	Pro	Pro	His	Сув	Phe 45	Lys	Arg	Ser	•	
15	* * * *	.•			:		1 1.	••			٠.	•		;	: .	• •		
		CCT															19:	2
	Ile	Pro	Arg	Ser	Phe	Ser	Tyr	Leu	Ile	\mathtt{Trp}	Asp	Ile	Ile	Ile	Ala	Ser		
	. : : * *	50				•	55					. 60	٠.		٠			
20		TTC															24	0
	Сув 65	Phe	тут	тут	vaı	70	Thr	Thr	ıyr	Pne	75	Leu	rein	PIO	uis	80		
											• • •							
		TCC															28	8
25	Leu	Ser	Tyr	Phe		Trp						Cys	Gln	Gly		Val		
					85		٠	:	•	90			٠:		95			
	СТА	ACC	GGC	GTC	TGG	GTC	ATA	GCC	CAC	GAG	TGC	GGC	CAC	CAC	GCC	TTC	33	6
		Thr																
30				100					105					110				
30		GAC		~~~	maa		~~	~~	3.00	ama	~~m	OTTO	3.000	TT C	CNC	TO C	38	 A
		Asp															36	*
	261	Asp	115	GIII	11D	Dea	ASP	120	****	V41	G. y		125	1	****	001		
35		CTC															43	2
	Phe	Leu	Leu	Val	Pro	Tyr		Ser	Trp	Lys	Tyr		His	Arg	Ser	His		
		130					135			٠		140						
																AAG	48	0
40	His	Ser	Asn	Thr	Gly	Ser	Leu	Glu	Arg	qzA	Glu	Val	Phe	Val	Pro			
	145					150					155					160		
	AAG	AAG	TCA	GAC	ATC	AAG	TGG	TAC	GGC	AAG	TAC	CAC	AAC	AAC	CCT	TTG	52	8
	Lys	Lys	Ser	qaA,	Ile	Lys	\mathtt{Trp}	Tyr	Gly	Lys	Tyr	His	Asn	Asn	Pro	Leu		
45					165					170					175			
	GGA	CGC	ACC	GTG	ATG	TTA	ACG	GTT	CAG	TTC	ACT	CTC	GGC	TGG	CCG	TTG	57	6
		Arg																
		_		180					185					190				
50	ም ክ ለ	TTA	CCC	Φ π∕	מממ	ርጥር	ጥርር	CCA	ልርያኦ	CCT	ጥልሮ	GAC	CCC	GGC	بانلىنلە _.	CCT	62	4
		Leu															02	•
	-7-	DC u	195	2110				200	3		-1-		205	1				
	* 1 *																	

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						_				CGT Arg			672
5										GGT Gly			720
10										TTC Phe 255			768
15								_		ACT Thr			816
		 								GAG Glu			864
20		 								GGA Gly	_		912
25										CAT His			960
30										AAG Lys 335			1008
										CCG Pro			1056
35										GAA Glu			1104
40										AAG Lys	TTA Leu	T	1153
	GA			. ::									1155
4 5	(2)	NFOR											

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 384 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

32

	(x)	L) :	SEQU	BNCE	DES	CRIP	rion	: S	EQ II	D NO	:8:					
5	Met 1	Gly	Ala	Gly	Gly 5	Arg	Met	Gln	Val	Ser 10	Pro	Pro	Ser	Lys	Lys 15	Ser
	Glu	Thr	Ąsp	Thr 20	Ile	Lys	Arg	Val	Pro 25	Сув	Glu	Thr	Pro	Pro 30	Phe	Thr
10	Val	Gly	Glu 35	Leu	Lys	Lys	Ala	Ile 40	Pro	Pro	His	Cys	Phe 45	ГÀ̀	Arg	Ser
	Ile	Pro 50	Arg	Ser	Phe	Ser	Tyr 55	Leu	Ile	Trp	Asp	Ile 60	Iļe	Ile	Ala	Ser
15	Суs 65	Phe	Tyr	Tyr	Val	Ala 70	Thr	Thr	Tyr	Phe	Pro 75		Leu	Pro	His	Pro 80
	Leu	Ser	Tyr	Phe	Ala 85	Trp	Pro	Leu	Tyr	Trp 90		Сув	Gln	Gly	Сув 95	Val
20	Leu	Thr	Gly	Val: 100	Trp	Val	Ile	Ala	His 105	Glu	Cys	Gly	His	His 110	Ala	Phe
25	Ser	Asp	Tyr 115	Gln	Trp	Leu	qaA	Asp 120	Thr	Val	Gly	Leu	Ile 125	Phe	His	Ser
	Phe	Leu 130	Leu	Val	Pro	Tyr	Phe 135	Ser	Trp	Lys	Tyr	Ser 140	His	Arg	Ser	His
30	His 145	Ser	Asn	Thr	Gly	Ser 150	Leu	Glu	Arg	Asp	Glu 155	Val	Phe	Val	Pro	Lys 160
	Lys	Lys	Ser	qaA	Ile 165	Lys	Trp	Tyr	Gly	Lys 170	Tyr	His	Asn	Asn	Pro 175	Leu
35	Gly	Arg	Thr	Val 180	Met	Leu	Thr	Val	Gln 185	Phe	Thr	Leu	Gly	Trp 190	Pro	Leu
	Tyr	Leu	Ala 195	Phe	Asn	Val	Ser	Gly 200	Arg	Pro	Tyr	Asp	Gly 205	Gly	Phe	Arg
40	Сув	His 210	Phe	His	Pro	Asn	Ala 215	Pro	Ile	Тут	Asn	Asp 220	Arg	Glu	Arg	Leu
45	Gln 225	Ile	Tyr	Ile	Ser	Asp 230	Ala	Gly	Ile	Leu	Ala 235	Val	Сув	Tyr	Gly	Leu 240
45	Phe	Arg	Tyr	Ala	Ala 245	Gly	Gln	Gly	Val	Ala 250	Ser	Met	Val	Cys	Phe 255	Tyr
50	Gly	Val	Pro	Leu 260	Leu	Ile	Val	Asn	Gly 265	Phe	Leu	Val	Leu	Ile 270	Thr	Tyr
	Leu	Gln	His 275	Thr	His		Ser	280	Pro	His	Tyr	Asp	Ser 285	Ser	Glu	Trp

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	Asp :	Trp 290	Phe	Arg	Gly	Ala	Leu 295	Ala	Thr	Val	Ąsp	Arg 300	Asp	Tyr	Gly	Ile		
5	Leu 2 305	Asn	Lys	Val	Phe	His 310	Asn	Ile	Thr	qzA	Thr 315	His	Val	Ala	His	His 320		٠
	Pro	Phe	Ser	Thr	Met 325	Pro	His	Tyr	His	Ala 330	Met	Glu	Ala	Thr	Lys 335	Ala		
10	Ile 1	Lys	Pro	Ile 340	Leu	Gly	Glu	Tyr	Tyr 345	Gln	Phe	Asp	Gly	Thr 350	Pro	Val		
	Val :	_	Ala 355	Met	Trp	Arg	Glu	Ala 360	Lys	Glu	Сув	Ile	Tyr 365	Val	Glu	Pro		
15	Asp	Arg . 370		Gly	Glu	Lys	L ув 375	Gly	Val	Phe.	Trp	Tyr 380	Asn	Asn	Lys	Leu		• • •
20	(2) (i)) s	EQUI	atio	CHAI	RACTI	ERIS	rics			. 7,						-	
25	(ii)	((B) (C) (D)	LENG TYPE STRA TOPO	: n MDEI LOGY	nucle ONESS (:]	eic a S: a Linea	cid ingl	.e	l							-	
30	(xi)			ence egtg(: S1	II QE) N O:	;9:							23
	(2) (i)) 5	EQU	MATI(ENCE	CHAI	RACT					·	••				•		
35		((A) (B) (C) (D)	STRA	eth: e: i andei ology	nucle ONESS	3: £	cid ingl										
40	(ii)) 1	40LE	CULE	TYP	E: 1	ANO.	(gen	omic)								
	(xi		-	ence rgct			TION	: S	BQ I	D NO	:10:							21
	(2)			MATI			EQ I	D NO	:11:									
4 5	(i			ENCE LENC		23	base	e pa										
50			(C) (D)	STR	ANDE	DNES:	S: :	sing: ar										
	(ii) 1	MOLE	CULE	TYP	E:	DNA	(gen	omic)	•							

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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
5	CATGGGTGCA GGTGGAAGAA TGC	23
5	(2) INFORMATION FOR SEQ ID NO:12:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	• •
15 ·	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
	TCTTTCACCA TCATCATATC C	21
00	(2) INFORMATION FOR SEQ ID NO:13:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	• .
25	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
<i>30</i>	GTCTGGGTCA TAGCCCACG	19
	(2) INFORMATION FOR SEQ ID NO:14:	
35	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
40	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
	GTCTGGGTCA TAGCCCACA	19
45	(2) INFORMATION FOR SEQ ID NO:15:	
50	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	

SEQUENCE DESCRIPTION: SEQ ID NO:15:

13 .

CTGGGTCATA GCCCATG 17 5 (2) INFORMATION FOR SEQ ID NO:16: (i) SEQUENCE CHARACTERISTICS: 17 base pairs · (A) LENGTH: 10 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear MOLECULE TYPE: DNA (genomic) (ii) 15 SEQUENCE DESCRIPTION: SEQ ID NO:16: (xi) 17 CTGGGTCATA GCCCACA 20 Claims 25 1. A method for altering fatty acid composition in plant seeds, comprising the steps of: a) introducing a recombinant nucleic acid construct into a plant, said construct comprising at least one seedspecific regulatory sequence operably linked in sense orientation to a mutant delta-12 fatty acid desaturase gene encoding a protein having a mutation in a His-(Asp/Glu)-Cys-(Gly/Ala)-His amino acid region; 30 b) obtaining progeny from said plant, said progeny producing seeds having said altered fatty acid composition; c) producing seeds having said altered fatty acid composition. 2. A method as claimed in claim 1, wherein said altered fatty acid composition comprises a decreased level of linoleic acid.

(xi)

3. A method as claimed in claim 2, wherein said altered fatty acid composition comprises from about 1.0% to about 10.0% linoleic acid, based on total fatty acid composition.

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4. A method as claimed in any one of claims 1 to 3, wherein said altered fatty acid composition comprises from about 69% to about 90% oleic acid, based on total fatty acid composition.

5. A method for altering fatty acid composition in plant seeds, comprising the steps of:

a) introducing a recombinant nucleic acid construct into a plant, said construct comprising at least one seedspecific regulatory sequence operably linked in sense orientation to a mutant delta-15 fatty acid desaturase gene encoding a protein having a mutation in a His-(Asp/Glu)-Cys-(Gly/Ala)-His amino acid region;

b) obtaining progeny from said plant, said progeny producing seeds having said altered fatty acid composition;

c) producing seeds having said altered fatty acid composition.

6. A method as claimed in claim 5, wherein said altered fatty acid composition comprises decreased levels of α-linolenic acid.

A method as claimed in any one of claims 1 to 6, wherein said construct comprises a full-length coding sequence of said mutant gene.

- 8. A method as claimed in any one of claims 1 to 7, wherein the plant is a Brassica canola plant.
- A method as claimed in any one of claims 1 to 8, wherein said mutant desaturase gene encodes a microsomal gene product.
- 10. A method as claimed in any one of claims 1 to 9, wherein said mutant desaturase gene encoding said protein has a non-conservative amino acid substitution in said region.
- 11. A method as claimed in claim 10, wherein said non-conservative amino acid substitution comprises the sequence 10 His-Lys-Cys-Gly-His.
 - 12. A method as claimed in any one of claims 1 to 11, wherein said mutant desaturase gene is derived from a <u>Brassica napus</u> plant.
- 13. A method for altering fatty acid composition in plant seeds wherein one or more recombinant nucleic acid constructs are introduced into said plant, said one or more constructs comprising:
 - a) at least one seed-specific regulatory sequence operably linked in sense orientation to a mutant delta-12 fatty acid desaturase gene encoding a protein having a mutation in a His-(Asp/Glu)-Cys-(Gly/Ala)-His amino acid region, as defined in any one of claims 1 or 7 to 12; and
 - b) at least one seed-specific regulatory sequence operably linked in sense orientation to a mutant delta-15 fatty acid desaturase gene encoding a protein having a mutation in a His-(Asp/Glu)-Cys-(Gly/Ala)-His amino acid region, as defined in any one of claims 2 or 7 to 12,
- 25 said mutant delta-12 and mutant delta-15 desaturase genes conferring altered fatty acid composition in seeds of said plant.
 - 14. A method as claimed in claim 13, wherein said altered fatty acid composition comprises from about 1.0% to about 10.0% linoleic acid and from about 1.0% to about 10.0% α -linolenic acid, based on total fatty acid composition.
 - 15. A plant obtainable by the method as defined in any one of claims 1 to 14.
 - 16. A plant cell obtainable by introduction of said recombinant nucleic acid construct as defined in any one of claims 1 to 14.
 - 17. A plant cell of a plant as defined in claim 15.
 - 18. A crushed seed produced by crushing the seeds obtainable by the method as claimed in any one of claims 1 to 14.
- 19. A vegetable oil extracted from the crushed seeds as defined in claim 18.
 - 20. A recombinant nucleic acid construct effective for altering fatty acid composition in seeds, wherein said construct is as defined in any one of claims 1 to 14.
- 21. A mutant delta-12 fatty acid desaturase comprising the amino acid sequence of SEQ ID NO:4 or SEQ ID NO:8 or a functionally equivalent variant, derivative or fragment thereof.
 - 22. A nucleic acid fragment encoding the mutant delta-12 fatty acid desaturase defined in claim 21.

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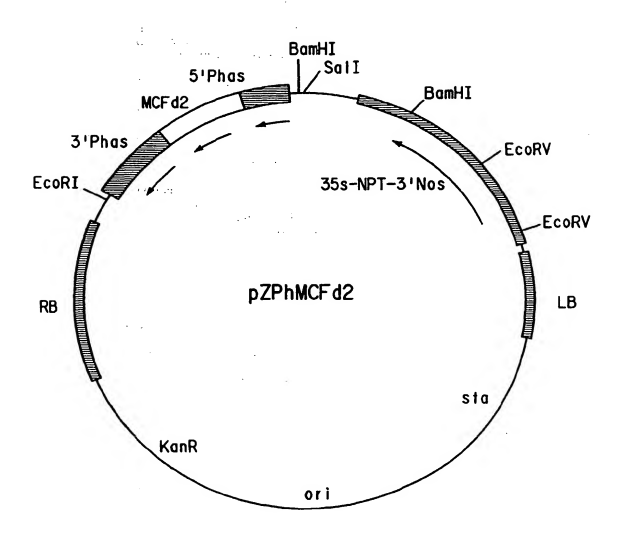


FIG.1

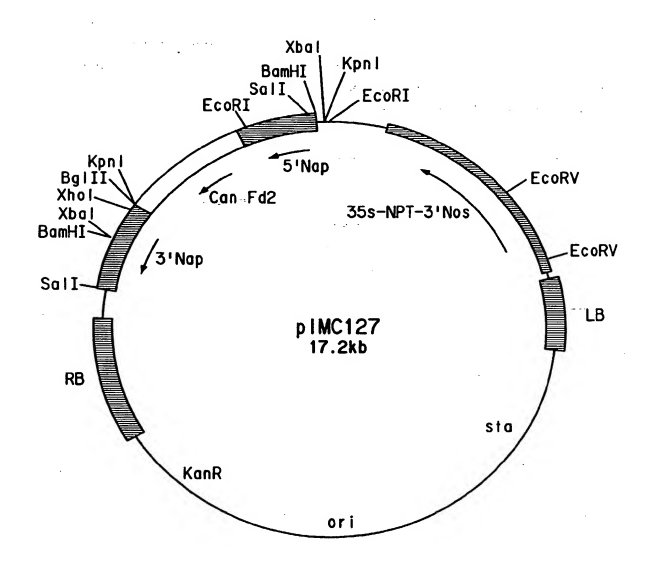
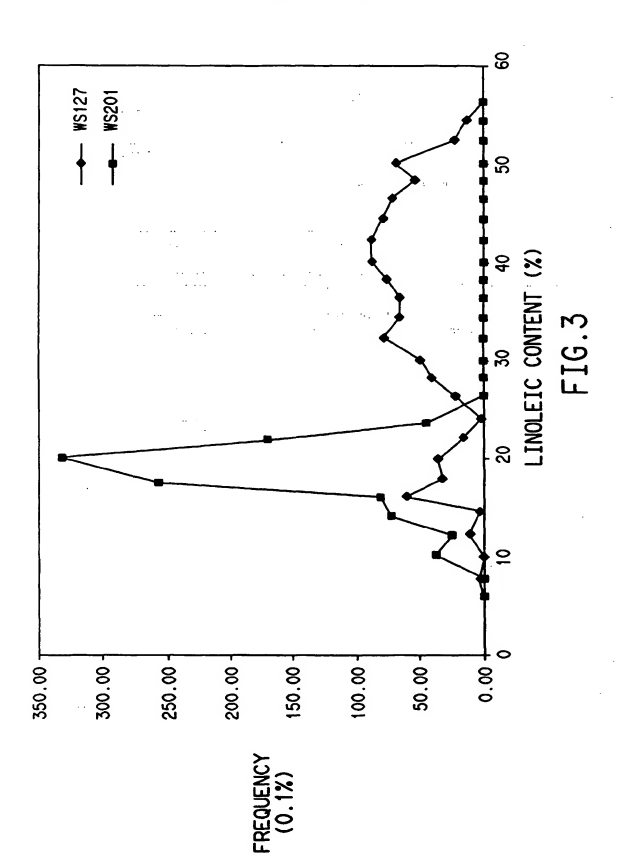
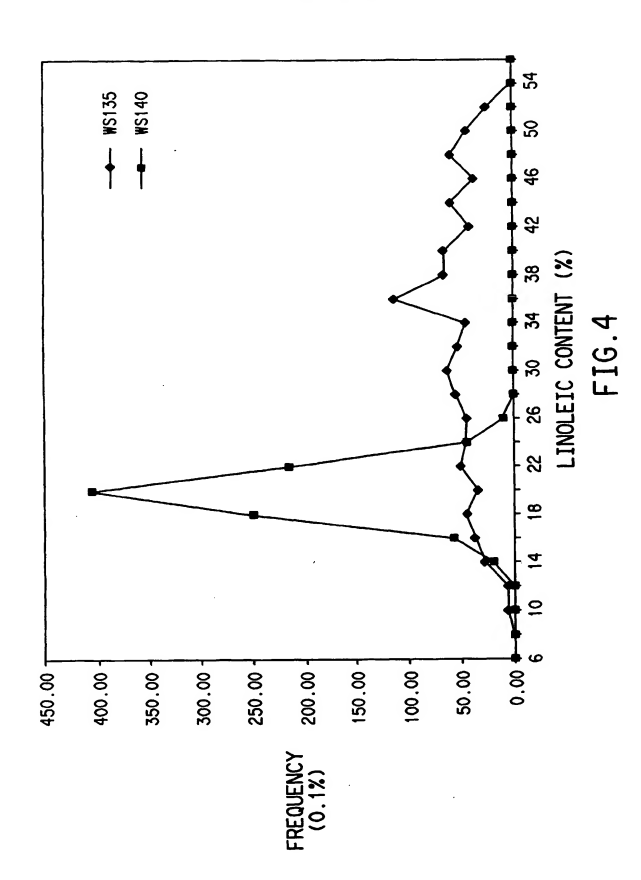


FIG.2





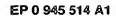


EUROPEAN SEARCH REPORT

Application Number EP 98 30 2322

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	The present search report has			
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